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(71) Applicants: BRESATEC LTD. [AU/AU]; P.O. Box 11, Rundie Mall, Adelaide, S.A. 5000 (AU). ST. VINCENT'S HOSPI-TAL (MELBOURNE) LIMITED [AU/AU]; 41 Victoria Parade, Fitzroy, VIC 3065 (AU).

(72) Inventors: PEARSE, Martin, J.; 37 Chute Street, Mordialloc, VIC 3195 (AU). CRAWFORD, Robert, J.; 7 Mirani Court, West Lakes Shore, S.A. 5020 (AU). ROBBINS, Allan, J.; Lot 8, Barker Road, Waterloo Comer, S.A. 5110 (AU). RATHJEN, Peter, D.; 1 Mimosa Avenue, Blackwood, S.A. 5051 (AU). d'APICE, Anthony, J., F.; 988 Burke Road, Balwyn, VIC 3103 (AU).

(74) Agent: MAXWELL, Peter, Francis; Peter Maxwell & Associates, Blaxland House, 5 Ross Street, North Parramatta, NSW 2151 (AU).

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(57) Abstract

Human pre-formed xenoantibodies play an important role in the hyperacute rejection response in human xenotransplantation. Disclosed are materials and methods for removing or neutralizing such antibodies. Also disclosed are materials and methods for reducing or eliminating the epitopes in the donor organs that are recognized by such antibodies. Such epitopes are formed as the result of activity by the enzyme α -1,3 galactosyltransferase. The porcine gene encoding α -1,3 galactosyltransferase is disclosed, as are materials and methods for inactivating ("knocking out") the α-1,3 galactosyltransferase gene in mammalian cells and embryos. Included are nucleic acid constructs useful for inactivating the α -1,3 galactosyltransferase gene in a target cell. Also disclosed is a novel leukemia inhibitory factor (T-LIF) that is useful for maintenance of embryonic stem cells and primordial germ cells in culture.

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MATERIALS AND METHODS FOR MANAGEMENT OF HYPERACUTE REJECTION IN HUMAN XENOTRANSPLANTATION

Field of the Invention

This invention relates generally to the field of xenotransplantation. In particular this invention relates to methods and materials for reduction or elimination of the hyperacute rejection response in humans. More particularly, this invention relates to methods for treating human serum to reduce or eliminate hyperacute rejection. This invention also relates to methods and materials for generating non-human organs lacking or having reduced α 1,3 galactosyl transferase activity.

15 Background of the Invention

It is widely acknowledged that there is an acute, worldwide shortage of human organs for transplantation. This is in spite of legislative changes and education programs to increase public awareness of the problem. In the United States, for example, there is an estimated annual shortfall of approximately 18,000 kidneys/year. Similarly, in Australia in 1992, only 41% of renal patients awaiting transplantation received transplants. In Japan the imbalance between supply and demand is even greater due to religious prohibitions on the use of organs from cadaveric donors.

The benefits of transplantation can be seen by comparing the rehabilitation rates of transplant patients with those of dialysis patients. In Australia and New Zealand, the majority of transplant patients (60%) are capable of full time work or school with a further 10% in part time work, while only 7% are unfit for work. In

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contrast, 23% of dialysis patients are capable of full time work or school, with 15% involved in part time work and 20% unfit for work. The remainder are "retired." Fifteenth Report of the Australia and New Zealand Dialysis and Transplant Registry (ANZDATA), Queen Elizabeth Hospital, Woodville, S.A., APS Disney, ed. (1992).

The direct financial cost of dialysis in Australia and New Zealand is approximately \$A45,000/patient/year.

10 In addition, indirect costs due to unemployment and sickness are higher in dialysis patients and the social costs are considerable. Transplantation engenders an expense of approximately \$A30,000/patient in the first year and \$A14,000/patient/year thereafter. These statistics indicate that a) transplantation is the optimal therapy for end stage renal failure; b) there is an undersupply of donor kidneys; and c) present strategies aimed at increasing the transplant rate have been less than successful. There are, in addition, serious shortages of other transplantable organs including hearts, livers, lungs and pancreases.

The use of xenografts (transplants between species) is one option for overcoming the short supply of human organs for transplantation. Non-viable, non25 antigenic xenografts are commonly used in vascular reconstruction (bovine arteries) and in cardiac surgery (porcine cardiac valves). However, despite their occasional use in the past, immunological barriers have prevented the common use of viable xenografts. Between 30 1964 and 1991 a total of 27 non-human primate to human organ xenografts was reported; the longest reported patient survival was 9 months. Two liver transplants from baboon to human were recently performed in anticipation that modern immunosuppressive therapies could cope with the severe rejection problems likely to

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occur in xenotransplantation. To date, the course of one of these patients has been reported, and in this case rejection was not the direct cause of death. Starzl et al., Baboon-to-Human Liver Transplantation. Lancet 341: 5 65-71 (1993). This clinical experience indicates that a) non-human organs can function and support human life; b) rejection episodes can be reversed by conventional anti-rejection therapy; and c) the mechanisms of rejection are similar, in principle, to those in allograft rejection.

It is unlikely that primates will be a satisfactory source of organs for xenotransplantation. Most are endangered species, breed slowly in the wild and poorly in captivity. The baboon is an exception to these generalizations, but other disadvantages limit the usefulness of this species. Baboons have single pregnancies, long gestation times, are difficult and expensive to maintain and may be infected with or carry organisms, particularly viruses, that are pathogenic in humans. For hearts and kidneys where organ size may be a consideration, the smaller primates are unsatisfactory as donors to human adults. Finally, the use of primates is likely to arouse considerable opposition from the public.

These difficulties have led to renewed interest in the use of non-primate species as organ donors for human patients. The pig is a widely acknowledged choice for xenotransplantation into humans. The pig erythrocyte diameter (6.5µm) and, by implication, its capillary size, are similar to humans, facilitating connection of xenografts to the human circulatory system. The pig breeds well in captivity, has a short gestation time and produces large litters. In addition, pigs can be bred and maintained in low pathogen facilities, can be reared to any size and do not arouse the level of public reaction associated with primates.

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The immunological barriers to use of pig organs in human patients include a) an immediate severe ("hyperacute") rejection phenomenon that develops in minutes to hours after transplantation, and b) a proposed 5 acute rejection that develops in days to weeks. Once the hyperacute rejection phenomenon has been overcome, it is expected that normal acute rejection would ensue. This form of rejection is thought to be similar to that experienced with allografts (transplants between 10 individuals of the same species) and should be amenable to normal immunosuppressive therapies.

Both preformed "natural antibodies"

(xenoantibodies) and complement regulating factors in human serum are thought to be involved in the process of hyperacute rejection. Hyperacute rejection is thought to be initiated when xenoantibodies bind to epitopes on the endothelium of a donor organ, activating the classical complement pathway.

Summary of the Invention

20 A purified and isolated nucleic acid molecule of the present invention comprises the porcine nucleic acid sequence depicted in Figure 4 (SEQ ID NO: 7), which encodes a porcine polypeptide having $\alpha-1,3$ galactosyltransferase activity. Variations on this 25 sequence that may be routinely generated by the skilled artisan include those sequences corresponding to Figure 4 but varying within the scope of the degeneracy of the genetic code. That is, the present invention includes variants of the sequence set out in Figure 4, readily 30 determined by the skilled artisan, that code for the same amino acid sequence encoded by the sequence set out in The present invention also includes a purified and isolated nucleic acid molecule that encodes a porcine α -1,3 galactosyltransferase and that hybridizes

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under standard high stringency conditions with a sequence complementary to the sequence set out in Figure 4, or with a sequence complementary to a variation of the sequence set out in Figure 4 within the scope of the degeneracy of the genetic code. The complementary strands to the above-described nucleic acid sequences are readily determined by standard methods, and are also within the scope of the present invention.

Within the parameters set out in the preceding 10 paragraph, the present invention includes variants of the porcine α -1,3 galactosyltransferase coding sequence that preserve the functional characteristics of the native gene product. Such variants include, for example, minor nucleotide variations in the 5' untranslated region or in 15 various coding regions of the disclosed sequence. Minor amino acid variations deriving from changes in the coding regions, that leave a functional $\alpha-1,3$ galactosyltransferase catalytic site, membrane anchor domain and stem region as described below, are within the 20 scope of the present invention. Such routine variations in nucleic acid and amino acid sequences can be identified by those having ordinary skill in the art based on the sequence and structural information provided herein.

As used herein, "high stringency conditions" are those hybridization conditions generally understood by the skilled artisan to reflect standard conditions of high stringency as set out in widely recognized protocols for nucleic acid hybridization. See, e.g., Sambrook et al, Molecular Cloning: A Laboratory Manual (2nd Edition), Cold Spring Harbor Laboratory Press (1989), pp. 1.101 - 1.104; 9.47 - 9.58 and 11.45 - 11.57. Generally, these conditions reflect at least one wash of the hybridization membrane in 0.05x to 0.5x SSC with 0.1% SDS at 65°C, or washing conditions of equivalent stringency.

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The present invention also includes a host cell transformed with any of the above-described purified and isolated nucleic acid molecules, as well as a porcine α-1,3 galactosyltransferase encoded by such transforming nucleic acid molecules and expressed from the host cell. Methods for transforming appropriate host cells and for expressing polypeptides from such host cells are known in the art and are described, for example, in Sambrook et al., (1984), pp. 12.2-12.44; 16.3-17.44.

The invention further includes a DNA construct useful for inactivating the porcine α-1,3 galactosyltransferase gene by insertion of a desired DNA sequence into an insertion site of the gene. As used herein, the term "α-1,3 galactosyltransferase gene" includes the exons encoding or potentially encoding α-1,3

galactosyltransferase, introns contiguous with such exons, and regulatory elements associated with such exons and introns. The DNA construct includes the desired DNA sequence flanked by first and second homology sequences.

These first and second homology sequences are sufficiently homologous, respectively, to first and second genomic sequences flanking the insertion site to allow for homologous recombination of the DNA construct with the porcine α -1,3 galactosyltransferase gene when

25 the DNA construct is introduced into a target cell containing the porcine α -1,3 galactosyltransferase gene. Preferably the insertion site is within exon 4, exon 7, exon 8 or exon 9 of the porcine α -1,3

galactosyltransferase gene. The desired DNA sequence is preferably a selectable marker, including but not limited to the neo^R gene, the hydromycin resistance (hyg^R) gene and the thymidine kinase gene. The desired DNA sequence may be bordered at both ends by FRT DNA elements, with stop codons for each of the three reading frames being

35 inserted 3' to the desired DNA sequence. Presence of the

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FRT elements allows the selectable marker to be deleted from the targeted cell, and the stop codons ensure that the α -1,3 galactosyltransferase gene remains inactivated following deletion of the selectable marker.

The invention further includes a DNA construct useful for inactivating the murine $\alpha-1,3$ galactosyltransferase gene by insertion of a desired DNA sequence into an insertion site of the gene. The DNA construct includes the desired DNA sequence flanked by 10 first and second homology sequences. These first and second homology sequences are sufficiently homologous, respectively, to first and second genomic sequences flanking the insertion site to allow for homologous recombination of the DNA construct with the murine $\alpha-1,3$ 15 galactosyltransferase gene when the DNA construct is introduced into a cell containing the murine $\alpha-1,3$ galactosyltransferase gene. Preferably the insertion site is within exon 4, exon 7, exon 8 or exon 9 of the murine α -1,3 galactosyltransferase gene. The desired 20 DNA sequence is preferably a selectable marker, including but not limited to the neo^R gene, the hyg^R gene and the thymidine kinase gene. The desired DNA sequence may be bordered at both ends by FRT DNA elements, with stop codons for each of the three reading frames being 25 inserted 3' to the desired DNA sequence. Presence of the FRT elements allows the selectable marker to be deleted from the targeted cell, and the stop codons ensure that the α -1,3 galactosyltransferase gene remains inactivated following deletion of the selectable marker.

The invention also includes methods for generating a mammalian totipotent cell having at least one inactivated (non-functional) α-1,3 galactosyltransferase allele, where the totipotent cell is derived from a mammalian species in which alleles for the α-1,3 galactosyltransferase gene normally are present and

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functional. A "functional" allele is capable of being transcribed and translated to produce a polypeptide having an activity the same as or substantially similar to the native $\alpha-1$,3 galactosyltransferase. The methods include providing a plurality of cells characterized as totipotent cells of the aforementioned mammalian species, introducing into the totipotent cells a nucleic acid construct effective for inactivating the $\alpha-1$,3 galactosyltransferase gene by insertion of a desired DNA sequence into an insertion site of the gene through homologous recombination, and then identifying a totipotent cell having at least one inactivated $\alpha-1$,3 galactosyltransferase allele.

The totipotent cells can include, without limitation, embryonic stem (ES) cells, primordial germ cells (PGC's) and eggs. The cells can be taken from a variety of mammalian species in which alleles for the α -1,3 galactosyltransferase gene are present and functional, including without limitation murine and porcine species.

The invention further includes methods for generating a mammal lacking a functional $\alpha-1,3$ galactosyltransferase gene, where the mammal belongs to a species having a functional α -1,3 galactosyltransferase The methods include providing a mammalian 25 gene. totipotent cell having at least one inactivated $\alpha-1,3$ galactosyltransferase allele, where the totipotent cell is derived from the aforementioned mammalian species having a functional α -1,3 galactosyltransferase gene, 30 manipulating the totipotent cell such that mitotic descendants of the cell constitute all or part of a developing embryo, allowing the embryo to develop to term, recovering a neonate individual derived from the embryo, and raising and breeding the neonate to obtain a 35 mammal homozygous for an inactivated $\alpha-1,3$

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galactosyltransferase alleles, i.e., a mammal in which both α -1,3 galactosyltransferase allele are inactivated.

The totipotent cells can include, without limitation, ES cells, PGC's and eggs. The cells can be 5 taken from a variety of mammalian species in which alleles for the α -1,3 galactosyltransferase gene are present and functional, including without limitation murine and porcine species. ES cells and PGC's are manipulated in various ways such that their mitotic 10 descendants are found in a developing embryo. manipulations can include, without limitation, injection into a blastocyst or morula, co-culture with a zona pellucida-disrupted morula, and fusion with an enucleated zygote. Cells injected into a blastocyst- or morula-15 stage embryo become incorporated into the inner cell mass of the blastocyst embryo, giving rise to various differentiated cell types of the resulting embryo, including in some cases germ cells. The embryo derived from such manipulations is a chimera composed of normal 20 embryonic cells as well as mitotic descendants of the introduced ES cells or PGC's. Alternatively, chimeric embryos can be obtained by co-culturing at least one ES cell or PGC with a morula embryo in which the zona pellucida is sufficiently disrupted to allow direct 25 contact between the ES cell/PGC and at least one cell of the morula. The zona pellucida-disrupted embryo may be an embryo that is completely free of the zona pellucida. Finally, the genome of an ES cell or PGC can be incorporated into an embryo by fusing the ES cell/PGC 30 with an enucleated zygote. Such a procedure is capable of generating a non-chimeric embryo, i.e., an embryo in which all nuclei are mitotic descendants of the fused ES cell/PGC nucleus. The resulting embryos are implanted in a recipient female, or surrogate mother, and allowed to 35 develop to term.

When eggs, as opposed to ES cells or PGC's, are directly injected with a nucleic acid construct effective for inactivating the α -1,3 galactosyltransferase gene, the eggs can be manipulated to form an embryo by implanting into a recipient female.

The invention also includes a mammal, produced through human intervention, that lacks a functional α -1,3 galactosyltransferase gene. The mammal belongs to a species in which the α -1,3 galactosyltransferase gene is normally present and functional. The mammal can be, without limitation, a mouse or a pig.

The invention further includes a purified and isolated nucleic acid molecule comprising a nucleic acid sequence selected from the group consisting of (1) the nucleic acid sequence depicted in Figure 26 (SEQ ID NO: 25), (2) a sequence corresponding to the sequence of (1) within the scope of the degeneracy of the genetic code, and (3) a sequence that encodes murine T-LIF and that hybridizes under standard high stringency conditions with a sequence complementary to the sequence of (1) or (2). The complementary strands to the above-described nucleic acid sequences are readily determined by standard methods, and are also within the scope of the present invention.

The present invention also includes a host cell transformed with any of the purified and isolated nucleic acid molecules described in the preceding paragraph, as well as a T-LIF polypeptide encoded by such transforming nucleic acid molecules and expressed from the host cell.

30 The invention further includes a purified and isolated nucleic acid molecule comprising a nucleic acid sequence selected from the group consisting of (1) the nucleic acid sequence depicted in Figure 27 (SEQ ID NO: 31), (2) a sequence corresponding to the sequence of (1) within the scope of the degeneracy of the genetic code,

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and (3) a sequence that encodes human T-LIF and that hybridizes under standard high stringency conditions with a sequence complementary to the sequence of (1) or (2). The complementary strands to the above-described nucleic acid sequences are readily determined by standard methods, and are also within the scope of the present invention.

The present invention also includes a host cell transformed with any of the purified and isolated nucleic acid molecules described in the preceding paragraph, as well as a T-LIF polypeptide encoded by such transforming nucleic acid molecules and expressed from the host cell.

The invention further includes a method for eliminating or reducing hyperacute rejection of non-15 primate mammalian cells by human serum, comprising adding, to the human serum, a physiologically acceptable amount of galactose or a saccharide in which the terminal carbohydrate is an α galactose linked at position 1, prior to exposure of the human serum to the non-primate The amount of galactose or saccharide added is sufficient to reduce or eliminate the hyperacute rejection response. The saccharide can be, without limitation, melibiose, galactose $\alpha 1-3$ galactose or stachyose. Alternatively, the human serum can be treated 25 so as to be substantially depleted of immunoglobulin, IgM antibodies, anti-GAL IgM and IgG antibodies, or anti-GAL IgM antibodies. The invention further includes affinitytreated human serum substantially free of anti-GAL antibodies or of anti-GAL IgM antibodies.

BRIEF DESCRIPTION OF THE DRAWINGS

FIGURE 1 is a graphical representation of fluorescence intensity following immunofluorescent staining of porcine aortic endothelial cells with anti-

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GAL antibody alone or with anti-GAL antibody that was preincubated with selected saccharides.

FIGURE 2 shows the results of an experiment in which lysis of porcine aortic endothelial cells by human 5 serum and by purified anti-GAL antibodies was determined using a 51CR release assay.

FIGURE 3 depicts physiograph tracings of perfused rat heart contractions in the presence of human serum with or without selected saccharides.

FIGURE 4 is a comparison of the porcine α-1,3 galactosyltransferase cDNA sequence with the corresponding murine and bovine sequences. PGTCD = porcine sequence. BOVGSTA = bovine sequence. MUSGLYTNG = murine sequence.

FIGURE 5 is a comparison of the porcine α -1,3 galactosyltransferase amino acid sequence with the corresponding murine and bovine amino acid sequences. PGT = porcine sequence. BGT = bovine sequence. MGT = murine sequence.

FIGURE 6 depicts the Sall restriction sites in four overlapping phage clones spanning a portion of the murine $\alpha-1,3$ galactosyltransferase genomic region.

FIGURE 7 is a detailed restriction map of murine α -1,3 galactosyltransferase subclone paGT-S5.5.

FIGURE 8 is a detailed restriction map of murine α -1,3 galactosyltransferase subclone paGT-S4.0.

FIGURE 9 is a detailed restriction map of murine $\alpha-1,3$ galactosyltransferase subclone pagt-S11.

FIGURE 10 is a detailed restriction map of murine 30 α -1,3 galactosyltransferase subclone paGT-S13.

FIGURE 11 is an additional detailed restriction map of murine α -1,3 galactosyltransferase subclone pagT-S5.5.

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FIGURE 12 is an additional detailed restriction map of murine α -1,3 galactosyltransferase subclone paGT-S4.0.

FIGURE 13 is a diagram of a knockout construct 5 carrying the 4.0 and 5.5kb Sal1 fragments from p α GT-S5.5 and p α GT-S4.0, which flank the Exon 9 Sal1 site.

FIGURE 14 depicts the 8.3kb and 6.4kb BglII fragments that are diagnostic for the uninterrupted α -1,3 galactosyltransferase gene and the targeted (inactivated) 10 α -1,3 galactosyltransferase gene, respectively, using the probes identified in the text.

FIGURE 15 is a schematic representation of the generation of a knockout construct using the vector p α GT-S5.5 as the starting vector.

15 FIGURE 16 sets out the nucleotide sequence of a neomycin resistance cassette used in the construction of a DNA construct for interrupting the α -1,3-GalT gene in mice.

FIGURE 17 is a diagram of one example of a final 20 knockout construct that has been sequenced to confirm the identity, copy number and orientation of the various inserts.

FIGURE 18 is a Southern blot of genomic DNA from various murine ES cell lines transformed with the 25 knockout construct of Figure 16, probed to reveal the diagnostic fragments depicted in Figure 14.

FIGURE 19 depicts the "long" PCR products derived from wild type and interrupted α -1,3-GalT genes using the designated primers.

FIGURE 20 is a Southern blot of long PCR products obtained from wild type and knockout mice.

FIGURE 21 depicts the PCR products used for identification of the interrupted (targeted) galT locus.

FIGURE 22 shows PCR products generated from mice 35 carrying interrupted (inactivated) GalT alleles.

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FIGURE 23 depicts the PCR products expected from PCR analysis of cDNA generated from α-1,3-GalT mRNA in normal and knockout mice. The ferrochelatase primers and PCR fragment represent a control demonstrating that cDNA synthesis had occurred.

FIGURE 24 shows the PCR fragments generated from cDNA obtained from RNA isolated from kidney (K), heart (H) and liver (L) of a wild-type mouse (+/+), a mouse heterozygous for the interrupted α-1,3-GalT allele (+/-) and a mouse homozygous for the interrupted α-1,3-GalT allele (-/-).

FIGURE 25 is a graphical representation of the relative protection of spleen cells, derived from GalT knockout mice, from lysis by human serum.

FIGURE 26 is a representation of the nucleotide sequence and deduced amino acid sequence for murine T-LIF.

FIGURE 27 is a representation of the nucleotide sequence and deduced amino acid sequence for human T-LIF.

FIGURE 28 is a Western blot of LIF polypeptides expressed from transfected COS cells.

FIGURE 29 is a diagram of the expression plasmid used for transfection of the COS cells of Figure 27.

FIGURE 30 is a Southern blot of PCR-amplified cDNA 25 from murine ES cells, using a LIF-specific probe.

DETAILED DESCRIPTION

Evidence presented herein establishes that a substantial portion of human pre-formed, anti-pig xenoantibodies recognize a specific terminal galactose linkage on the surface of pig endothelial cells. As demonstrated in experiments carried out by the present inventors, it is possible to reduce the titers of preformed xenoantibodies by adsorption with immobilized antigens containing the appropriate epitopes. This leads

to reduction or elimination of cellular responses associated with the hyperacute rejection response. Conversely, it is demonstrated to be possible to neutralize such antibodies by addition of appropriate carbohydrate antigens to human serum. In demonstrating the usefulness of these approaches, it was necessary to identify the relevant carbohydrate moieties and to demonstrate their efficacy in cultured cell systems and, importantly, in whole organs. As such, one approach to reducing or eliminating the hyperacute rejection response is identified as treatment of the recipient by eliminating or neutralizing the relevant antibody populations.

An alternative approach to xenotransplantation

15 would be elimination of the relevant epitope(s) in the
donor organ. This could be accomplished, for example, by
reducing or eliminating expression of the gene(s)
encoding the metabolic machinery responsible for
formation of the epitopes. The epitope defined by the α
20 1,3 galactose linkage (termed the GAL epitope) is
generated by the enzyme UDP-galactose:β-D-galactosyl-1,4N-acetyl-D-glucosaminide α-1,3 galactosyl- transferase
("α-1,3 galactosyltransferase" or "α-1,3-GalT"). This
enzyme transfers galactose to the terminal galactose

25 residue of N-acetyllactosamine-type carbohydrate chains
and lactosaminoglycans. The reaction catalyzed by α-1,3GalT may be summarized as follows:

UDP-Gal + Gal β -1,4-GlcNAc-R \Rightarrow Gal α -1,3-Gal β -1,4-GlcNAc-R + UDP

The α-1,3-Gal T enzyme is found in most mammals, but is not present in Old World monkeys and humans. For purposes of xenotransplantation, it is significant that humans and Old World monkeys have naturally occurring xenoantibodies directed against the GAL epitope. The use of pig organs lacking the GAL epitope could reduce or

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eliminate the hyperacute rejection of such organs by human recipients. The utility of such an approach is buttressed by the present inventors' demonstration that the GAL epitope is, in fact, central to the hyperacute rejection phenomenon in cells and whole organs. One approach to obtaining such organs would be to generate pigs in which the gene encoding the α-1,3-GalT enzyme is "knocked out" by homologous recombination.

Role of the GAL Epitope in Hyperacute Rejection

The present inventors have affinity purified 10 antibodies directed against the GAL epitope (anti-GAL antibodies) from human serum. This was accomplished with affinity columns comprising the appropriate epitopes (e.g., galactosyl-galactose or melibiose) attached to a 15 solid phase. Total anti-GAL IgG and IgM were obtained in one set of experiments. In an alternative approach, anti-GAL IgG was obtained by passage of serum over an affinity column with specificity for all proteins except albumin and IgG. The wash-through from this column was 20 then applied to a galactosyl-galactose affinity column and purified anti-GAL IgG was collected as the eluate. The obtained anti-GAL IgG can be further purified by passage over a protein G column, which specifically binds IgG but not other antibody isotypes. Conversely, the 25 wash-through from the above-described columns can be used as sources of total anti-GAL (IgG + IgM)-depleted serum or of anti-GAL IgG-depleted serum in further experiments. Preferably, the anti-GAL antibody preparations are characterized for protein content, molecular weight and 30 purity, and for antibody class and isotype.

To demonstrate the role of the GAL epitope in the hyperacute rejection response, it is necessary, first, to establish that IgG and IgM anti-GAL antibodies react with porcine cells and tissues. The present inventors

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investigated the binding of human anti-GAL antibodies to porcine cells and tissues using immunofluorescent staining. In this technique, selected human antibody preparations are reacted with intact porcine cells and then reacted with signal antibody comprising non-human anti-human IgG or IgM labeled with fluorescein isothiocyanate (FITC). Stained cells may be detected and quantified with a fluorescence-activated cell sorter (FACS) or other appropriate detection means. Other methods for detecting the presence of a bound antibody on a cell surface, for example through use of enzyme-labeled signal antibody reagents, are known to the skilled artisan.

Total anti-GAL (IgM and IgG), as well as 15 purified anti-GAL IgG, stained cells from a porcine epithelial cell line (PK1) as well as cells from a porcine aortic endothelial cell line (PAE). Neither anti-GAL (total IgM + IgG) antibody-depleted serum nor anti-GAL IgG-depleted serum gave detectable staining. 20 further investigate the specificity of the response, it is desirable to determine whether or not reactivity of the antibodies with porcine cells can be diminished or eliminated by prior exposure to one or more molecules suspected of comprising the epitope(s) in question. 25 this regard, the present inventors have established that antibody binding is inhibited by galactose and by disaccharides having terminal galactose residues in the al configuration. Staining was not inhibited with sugars having a terminal galactose in a $\beta 1\rightarrow 4$ configuration. 30 These results demonstrate the specificity of the antibody binding and the ability of appropriate sugars to inhibit such binding.

Reactivity of anti-GAL antibodies with cultured pig cells was confirmed using tissue sections of pig organs. Again, using a fluorescent signal antibody

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system, staining was seen with total anti-GAL IgM + IgG and with purified anti-GAL IgG but not with the anti-GAL antibody-depleted sera. Staining was particularly strong with kidney, heart and liver endothelium, with heart endocardium and with bile duct epithelium. The tissue binding was inhibited with melibiose but was not inhibited by other disaccharides not representative of the GAL epitope.

These data clearly indicate that the GAL epitope
is expressed at high levels on the endothelial cells of
arteries, veins and capillaries of porcine kidney, heart
and liver. In a xenograft situation, the endothelial
cells of these vessels come into direct contact with the
anti-GAL antibodies in human serum. The above results
are consistent with evidence that binding of these
antibodies (with attendant complement activation) is a
key component of the hyperacute rejection response.

To further investigate the specificities of naturally occurring xenoantibodies in human serum

20 directed against porcine antigens, the ability of human serum to cause agglutination of pig red blood cells was investigated. These studies revealed the presence of high levels of such antibodies in human serum. Moreover, sugars such as melibiose, stachyose, galactose and

25 fucose, having terminal residues in the α1-6 configuration, were found to inhibit agglutination in the μM to mM range. Sugars with other configurations were only inhibitory at very high doses, where the observed effects are likely due to simple changes in osmolarity or other non-specific mechanisms.

The above investigations establish a potential role for naturally occurring, human anti-GAL xenoantibodies in the complement-mediated destruction underlying hyperacute rejection. However, it is preferable to directly examine complement-mediated

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destruction of porcine cells in order to confirm the specificity of the GAL epitope and of anti-GAL antibodies in the process of lysis. To this end, the present inventors have examined the ability of human serum to 5 cause lysis of porcine cells.

To investigate complement-mediated destruction of cells, it is necessary to employ one or more assays that provide quantitative data on cell lysis. Preferably, such assays measure a cell-sequestered component that is released into the medium upon complement-mediated cell lysis. Such experiments should control for involvement of complement in the induced lysis by employing both native complement proteins as well as heat-inactivated complement. The present inventors have used a 51Cr-15 release assay and a lactate dehydrogenase (LDH)-release assay to investigate the complement-mediated lysis of porcine epithelial and endothelial cells by human serum.

In the ⁵¹Cr-release assay, porcine cells were pre-labeled with ⁵¹Cr and then incubated in the presence of heat-inactivated human serum plus rabbit complement (PAE's) or human complement in non-heat-inactivated normal human serum (PK₁'s). Release of ⁵¹Cr into the medium was measured with a gamma counter following addition of scintillation fluid. In the LDH-release assay, cells were labeled with LDH as per the manufacturer's instructions (Promega, USA). Release of LDH into the medium was measured using an ELISA format, with absorbance read at 492nm. For both assays, the ability of various sugars to inhibit the complementinduced lysis was also tested.

Similar results were obtained with the two unrelated porcine cell lines, PAE and PK₁, using both types of assays. The results clearly demonstrate that naturally occurring xenoantibodies (NXAb's) are responsible for initiating the complement-induced lysis

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of porcine cells. The present inventors have also established that IgM and not IgG antibodies are responsible for the lysis in this system. Moreover, heat inactivation of the complement preparations prevented lysis, providing further evidence that lysis of the porcine cells is a complement-dependent phenomenon. The present inventors have also shown that melibiose, but not lactose, protects the porcine cells from lysis. Importantly, the concentrations of sugar found to be effective in these studies covered the physiological range of blood sugar, i.e., about 10mM.

These results indicate that the anti-GAL NXAb's in normal human serum are primarily responsible for lysis of the porcine cells. As such, the binding of anti-GAL

NXAb's to the endothelial cells lining the blood vessels of a porcine xenograft, with attendant activation of the complement cascade, is likely to be a key component of the hyperacute rejection of porcine xenografts. This would also be the case with organs from other discordant species, such as rodents, sheep, cows and goats, all of which have active α-1,3-GalT genes in their genomes.

These conclusions are further supported in a whole-organ study performed by the present inventors. For this study, isolated and perfused rat hearts were used to further demonstrate the involvement of anti-GAL xenoantibodies in hyperacute rejection. Rat hearts were connected to a Langendorf perfusion apparatus, as described in Doring and Dehnert, The Isolated Perfused Heart According to Langendorf, Bionesstechnik-Verlag March GmbH, D7806, West Germany. The connected hearts were then stabilized by perfusion with a physiological buffer system, and perfused with the same buffer containing either melibiose or lactose (10mM). Human plasma was then added to a final concentration of 13% and

the effect of the added sugar on heart rate, strength of contraction and output were measured.

These results demonstrate in a whole-organ system that:

- 5 1) Perfusion with unmodified human plasma causes rapid loss of function.
- 2) Perfusion of a rat heart with human plasma in the presence of melibiose, which competes for binding with the anti-GAL antibodies, prolongs heart survival and 10 output. Lactose, however, which does not compete for binding with the anti-GAL antibodies, does not prolong heart survival.
- 3) Perfusion of a rat heart with anti-GAL antibody-depleted plasma prolongs heart survival and 15 output.
 - 4) If purified anti-GAL antibodies are added back to anti-GAL antibody-depleted plasma, the heart rapidly loses function

The present inventors' experiments with cultured cells, tissues and whole organs provide important confirmation that anti-GAL antibodies are a critical element in the hyperacute rejection response. Moreover, the disclosed results point to various approaches that can be employed to eliminate or reduce the hyperacute rejection of xenogeneic mammalian organs by humans.

For example, the intravenous administration of the specific disaccharide galactose α 1-3 galactose will block the naturally occurring anti-GAL antibodies of all classes and prevent them binding to their specific 30 epitopes on the surface of the endothelial cells of the xenograft, thus preventing them from initiating or participating in hyperacute rejection. The present inventors' results indicate that the concentration of galactose α 1-3 galactose required to achieve this effect

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is in a physiologically tolerated range. The experiments also indicate that various other carbohydrates can be substituted for the specific disaccharide. These include the monosaccharide galactose and various other di-, tri- or tetra-saccharides in which there is a terminal a galactose linked to the next sugar via position 1. These other sugars include, but are not limited to, melibiose and stachyose.

Likewise, prior to xenotransplantation, all or a

10 substantial portion of total IgM (that is, IgM of all
specificities) can be removed from the serum of the
patient by extracorporeal immunoabsorption.
Alternatively, anti-GAL antibodies of all classes can be
removed by extracorporeal immunoabsorption. Most

15 preferably, the patient's pre-formed natural anti-GAL IgM
antibodies can be removed. In this way, many or most of
the primary immunological agents of the hyperacute
response are eliminated, resulting in reduction or
elimination of the response following

20 xenotransplantation.

The α -1,3-GalT Gene as a Target for Suppressing the GAL Epitope

The present inventors have succeeded in cloning the entire coding region of the porcine α-1,3-GalT gene.

25 This is desirable for full exploitation of the gene in genetic engineering of pigs for purposes of human xenotransplantation. Previous attempts to obtain the entire coding region of the porcine gene have, to the knowledge of the inventors, failed to generate the 5'

30 coding regions. See, e.g., Dabkowski et al., Transplant. Proc. 25: 2921 (1993). The present inventors have employed a PCR-based approach to generate the full sequence. In designing the primers and experimental conditions required to obtain the 5' and 3' regions of

the gene, the present inventors overcame significant theoretical and practical obstacles to success.

Primers were selected on the basis of careful analysis of published sequences for the murine, bovine 5 and human α -1,3-GalT genes, the only published sequences available for this purpose. The present inventors' analysis revealed that in the reported sequence of the bovine cDNA, exon 3 (which is in the 5'-untranslated region) is missing. This had not been reported in the 10 literature. Thus, in order to find appropriate regions for deriving useful primer sequences, the mouse and bovine sequences had to be realigned. Even with the appropriate realignment, however, only one island of about 20 base pairs (bp) in the 5' untranslated region 15 displayed the desired homology (19 out of 20 bp) for design of a PCR oligonucleotide. The fact that the 5' untranslated regions of the mouse and bovine genes do not seem substantially related even upon optimal alignment would not be considered unusual by the ordinary skilled 20 artisan. This is because the 5' untranslated regions are often not well conserved between species. As such, the natural inclination would be to perform a less-thanexhaustive analysis and to conclude that design of PCR oligonucleotides based on homology from this region was 25 unlikely to be successful.

In the downstream 3'-untranslated region, the homology is less than obvious again. Various insertions and deletions had to be made in order to obtain proper alignment of the mouse and bovine sequences. Moreover, 30 to obtain a region of appropriate homology for design of PCR oligonucleotides, it was necessary to select a region approximately 200 bp downstream of the stop codon. Finally, to get the 5' and 3' primers to work properly, the present inventors found it necessary to drop the 35 annealing temperature by 9°C. These technical and

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theoretical hurdles to successful use of a PCR-based approach were overcome by the present inventors and allowed the entire coding sequence to be determined.

Analysis of the nucleotide sequence indicates that 5 a counterpart to murine exon 3 in the 5' untranslated region is not found in the porcine gene. The porcine sequence is similar to the bovine sequence in this regard. Analysis of the amino acid sequence demonstrates that the structure of the porcine $\alpha-1,3$ -GalT is similar 10 to that of other glycosyltransferases, and in particular is closely related to bovine and murine $\alpha-1,3$ -GalTs. each of these enzymes a short cytoplasmic amino-terminal domain of about 6 residues precedes a hydrophobic membrane-anchoring domain (extending from residues 7 to 15 22). The stem region, which serves as a flexible tether, and the catalytic domain, which catalyses the synthesis of α -1,3-GAL linkages, are located in the lumen of the Golgi and extend from amino acid 23 to the carboxyl terminus at amino acid 371. The precise boundary between 20 the stem and catalytic domains is not well-defined. Based on the suggested characteristics of the stem region, it appears to be the least conserved region and is rich in glycine and proline residues. Paulson and Colley, J. Biol. Chem. 264: 17615 (1989); Joziasse et 25 al., J. Biol. Chem. 267: 5534 (1992). The stem/catalytic boundary may occur around amino acid 60.

To generate constructs for inactivating genes by homologous recombination, the gene is preferably interrupted within an appropriate coding exon by insertion of an additional DNA fragment. Upon analysis of the full-length porcine nucleic acid sequence, the present inventors have identified exons 4, 7, 8 and 9 as preferred locations for disruption of the gene by homologous recombination. However, identification of these exons as preferred sites should not be construed as

limiting the scope of the present invention, as interruptions in exons 5 and 6 may be useful in particular cell types or in situations where less-than-complete inhibition of α -1,3-GalT gene expression is desired. Moreover, regulatory elements associated with the coding sequence may also present useful targets for inactivation.

In a preferred embodiment, a Sall site located within exon 9 of the mouse α-1,3-GalT gene at codons 22110 222 is chosen as the site for disruption of the murine coding sequence. For disruption of the porcine sequence, it is noted that the amino acids encoded by the corresponding porcine nucleotides are conserved, although the Sall site is not. In a preferred embodiment for 15 inactivation of the porcine gene, a Sall site is engineered into the corresponding location of the pig sequence for convenient construction of a knockout sequence. Sall cuts only rarely in genomic DNA. Since multiple restriction sites can be a problem in 20 manipulating large fragments of DNA, the presence of a Sall site in the exon is very useful since it is not likely that other Sall sites will be present at other

A gene coding for a selectable marker is generally used to interrupt the targeted exon site by homologous recombination. Preferably, the selectable marker is flanked by sequences homologous to the sequences flanking the desired insertion site. Thomas and Capecchi, Cell 51: 503-12 (1987); Capecchi, Trends in Genetics 5: 70-76 (1989). It is not necessary for the flanking sequences to be immediately adjacent to the desired insertion site. The gene imparting resistance to the antibiotic G418 (a neomycin derivative) frequently is used, although other antibiotic resistance markers (e.g., hygromycin) also may be employed. Other selection systems include negative-

locations in the knockout constructs.

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selection markers such as the thymidine kinase (TK) gene from herpes simplex. Any selectable marker suitable for inclusion in a knockout vector is within the scope of the present invention.

However, it is possible that in some circumstances it will not be desirable to have an expressed antibiotic resistance gene incorporated into the cells of a transplanted organ. Therefore, in a preferred embodiment, one or more genetic elements are included in the knockout construct that permit the antibiotic resistance gene to be excised once the construct has undergone homologous recombination with the α-1,3-GalT gene.

The FLP/FRT recombinase system from yeast 15 represents one such set of genetic elements. O'Gorman et al., Science 251, 1351-1355 (1991). FLP recombinase is a protein of approximately 45 kD molecular weight. is encoded by the FLP gene of the 2 micron plasmid of the yeast Saccharomyces cerevisiae. The protein acts by 20 binding to the FLP Recombinase Target site, or FRT; the core region of the FRT is a DNA sequence of approximately 34 bp. FLP can mediate several kinds of recombination reactions including excision, insertion and inversion, depending on the relative orientations of flanking FRT 25 sites. If a region of DNA is flanked by direct repeats of the FRT, FLP will act to excise the intervening DNA, leaving only a single FRT. FLP has been shown to function in a wide range of systems, including in the cultured mammalian cell lines CV-1 and F9, O'Gorman et 30 al., Science 251: 1351 (1991), and in mouse ES cells, Jung et al., Science 259: 984 (1993).

Targeted cells carrying a genomic copy of an antibiotic resistance gene flanked by direct repeats of the FRT are supplied with FLP recombinase by 1)

35 introduction into cells of partially purified FLP protein

by electroporation, or 2) transfection with expression plasmids containing the FLP gene. In this way, the antibiotic resistance gene is deleted by action of the FLP recombinase, and cells are generated that contain the inactivated α-1,3-GalT gene and are free of the exogenous antibiotic resistance gene.

Due to the relative infrequency of homologous recombination in targeted cells, most such cells will carry only one inactivated allele of the target gene. 10 That is rathe great majority of cells taken through a single round of transformation with an appropriate knockout construct will be heterozygotes. As used herein, the term "transformed" is defined as introduction of exogenous DNA into the target cell by any means known 15 to the skilled artisan. These methods of introduction can include, without limitation, transfection, microinjection, infection (with, for example, retroviralbased vectors), electroporation and microballistics. term "transformed," unless otherwise indicated, is not 20 intended herein to indicate alterations in cell behavior and growth patterns accompanying immortalization, density-independent growth, malignant transformation or similar acquired states in culture.

Although heterozygous cells can be used in the
25 methods of the present invention, various manipulations
can be employed to generate homozygous cells in culture.
For example, homozygous cells can be generated by
performing a second homologous recombination procedure on
cells heterozygous for the inactivated allele. If the
30 knockout construct used in the initial transformation
carried the neo^R gene, a second construct may be employed
in a second round of transformation in which the neo^R
gene is replaced with a gene conferring resistance to a
separate antibiotic (e.g., hygromycin). Cells resistant
35 to both G418 and hygromycin can be screened by Southern

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blots in order to detect any "double knockouts" (i.e., homozygotes). Both antibiotic resistance genes can be removed subsequently in a single procedure using FLP recombinase. By maintaining selection with G418, this approach ensures that the second construct does not simply replace the previously knocked-out allele, leaving the cells heterozygous.

Alternatively, the neo^R gene can be deleted from an original heterozygous cell using FLP recombinase and a second knockout procedure conducted using the original neo^R gene-containing construct. Double knockouts could be detected by Southern analysis. The newly introduced neo^R gene then could be deleted by FLP recombinase. This alternative approach does not allow one to direct the knockout construct specifically to the non-inactivated allele. Nevertheless, screening of appropriate numbers of targeted cells can lead to identification of cells homozygous for the inactivated locus.

Cellular Vehicles for Incorporation of Knockout Constructs 20 To create animals having a particular gene inactivated in all cells, it is necessary to introduce a knockout construct into the germ cells (sperm or eggs, i.e., the "germ line") of the desired species. Genes or other DNA sequences can be introduced into the pronuclei 25 of fertilized eggs by microinjection. Following pronuclear fusion, the developing embryo may carry the introduced gene in all its somatic and germ cells since the zygote is the mitotic progenitor of all cells in the Since targeted insertion of a knockout construct 30 is a relatively rare event, it is desirable to generate and screen a large number of animals when employing such an approach. Because of this, it can be advantageous to work with the large cell populations and selection criteria that are characteristic of cultured cell

systems. However, for production of knockout animals from an initial population of cultured cells, it is necessary that a cultured cell containing the desired knockout construct be capable of generating a whole animal. This is generally accomplished by placing the cell into a developing embryo environment of some sort.

cell into a developing embryo environment of some sort. Cells capable of giving rise to at least several differentiated cell types are hereinafter termed and a "pluripotent" cells. Pluripotent cells capable of giving 10 rise to all cell types of an embryo, including germ cells, are hereinafter termed "totipotent" cells. Totipotent murine cell lines (embryonic stem, or "ES" cells) have been isolated by culture of cells derived from very young embryos (blastocysts). Such cells are ..15 capable, upon incorporation into an embryo, of differentiating into all cell types, including germ cells, and can be employed to generate animals lacking a functional $\alpha-1,3$ -GalT gene. That is, cultured ES cells can be transformed with a knockout construct and cells 20 selected in which the $\alpha-1,3$ -GalT gene is inactivated through insertion of the construct within, for example, an appropriate exon. In fact, ES cell lines have been derived for both mice and pigs. See, e.g., Robertson, Embryo-Derived Stem Cell Lines. In: Teratocarcinomas and 25 Embryonic Stem Cells: A Practical Approach (E.J. Robertson, ed.), IRL Press, Oxford (1987); PCT Publication No. WO/90/03432; PCT Publication No. 94/26884. Generally these cells lines must be propagated in a medium containing a differentiation-inhibiting 30 factor (DIF) to prevent spontaneous differentiation and loss of mitotic capability. Leukemia Inhibitory Factor (LIF) is particularly useful as a DIF. Other DIF's useful for prevention of ES cell differentiation include, without limitation, Oncostatin M (Gearing and Bruce, The

35 New Biologist 4: 61-65 (1992); personal communication

from A. Smith), interleukin 6 (IL-6) with soluble IL-6 receptor (sIL-6R) (Taga et al., Cell 58: 573-81 (1989); personal communication from A. Smith), and ciliary neurotropic factor (CNTF) (Conover et al., Development 19: 559-65 (1993). Other known cytokines may also function as appropriate DIF's, alone or in combination with other DIF's.

As a useful advance in maintenance of ES cells in an undifferentiated state, the present inventors have identified a novel variant of LIF. In contrast to the previously identified forms of LIF which are extracellular, this new form of LIF (hereinafter T-LIF) is intracellularly localized. The transcript was cloned from murine ES cells using the RACE technique, Frohman et al., Proc. Natl. Acad. Sci. USA 85: 8998-9002 (1988), and subjected to sequence analysis. Analysis of the obtained nucleic acid sequence and deduced amino acid sequence indicates that T-LIF is a truncated form of the LIF sequence previously reported in the literature.

20 Expression of the T-LIF nucleic acid in an appropriate host cell yields a 17 kD protein that is unglycosylated. This protein is useful for inhibiting differentiation of murine ES cells in culture. The protein is expected to have a similar activity with porcine cells, since murine 25 D-LIF is effective at inhibiting both murine and porcine ES cell differentiation. The present inventors have also determined the sequence of the human form of T-LIF.

To generate a knockout animal, ES cells having at least one inactivated α-1,3-GalT allele are identified

30 and incorporated into a developing embryo. This can be accomplished through injection into the blastocyst cavity of a murine blastocyst-stage embryo, by injection into a morula-stage embryo, by co-culture of ES cells with a morula-stage embryo, or through fusion of the ES cell

35 with an enucleated zygote. The resulting embryo is

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raised to sexual maturity and bred in order to obtain animals, all of whose cells (including germ cells) carry the inactivated α-1,3-GalT allele. If the original ES cell was heterozygous for the inactivated α-1,3-GalT allele, several of these animals must be bred with each other in order to generate animals homozygous for the inactivated allele.

Although direct microinjection of DNA into eggs does not generate the large numbers of recombination

10 events obtained through transfecting large numbers of cultured cells, nevertheless direct injection of eggs can be a useful approach since this avoids the special manipulations (see above) required to turn a cultured cell into an animal. This is because fertilized eggs are, of course, quintessentially "totipotent" - i.e., capable of developing into an adult without further substantive manipulation other than implantation into a surrogate mother. To enhance the probability of homologous

recombination when eggs are directly injected with knockout constructs, it is useful to incorporate at least about 8 kb of homologous DNA into the targeting construct. In addition, it is also useful to prepare the knockout constructs from isogenic DNA. For example, for injection of porcine eggs, it is useful to prepare the constructs from DNA isolated from the boar whose sperm are employed to fertilize the eggs used for injection.

Embryos derived from microinjected eggs can be screened for homologous recombination events in several 30 ways. For example, if the GalT gene is interrupted by a coding region that produces a detectable (e.g., fluorescent) gene product, then the injected eggs are cultured to the blastocyst stage and analyzed for presence of the indicator polypeptide. Embryos with fluorescing cells, for example, are then implanted into a

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surrogate mother and allowed to develop to term.

Alternatively, injected eggs are allowed to develop and the resulting piglets analyzed by polymerase chain reaction (PCR) or reverse transcription PCR (RT/PCR) for evidence of homologous recombination.

Characterization of Knockout Animals

Animals having either one (heterozygous) or two (homozygous) inactivated GalT genes are characterized to confirm the expected alterations in gene expression and 10 phenotypic effect. For example, GalT mRNA should be absent from homozygous knockout animals. This can be confirmed, for example, with reverse transcription PCR (RT-PCR) using appropriate GalT-specific primers. addition, various tests can be performed to evaluate 15 expression of the GAL epitope in homozygous knockout animals. For example, anti-GAL antibodies and IB4 Lectin (which has an exclusive affinity for terminal α -Dgalactosyl residues) can be used in various assay or immunohistological formats to test for the presence of 20 the GAL epitope in an array of tissues. As another indication of GAL epitope status, lysis of cells by human serum can be tested through use of a 51chromium release assay.

EXAMPLE 1

- Affinity Purification of Human Anti-GAL Antibodies

 Anti-GAL antibodies were purified from normal heat
 inactivated AB serum (from CS1, Parkville, Victoria,
 Australia) using the following sets of procedures.
- A. <u>Preparation of total anti-GAL (IgG+IgM) antibodies</u>
 30 The following procedures are performed at 4°C.
 - 1. Desalt 15-30ml serum (in 3ml batches) by passage through a pre-equilibrated (20ml application buffer: 20mM $\rm K_2HPO_4$, 30mM NaCl, pH 8) Econo Pac 10DG (Bio-Rad, Richmond, USA) column. Alternatively, for large scale

preparations, desalt by dialysis exhaustively against application buffer.

- 2. Wash column with 4ml aliquots of application buffer. Collect and pool column eluates.
- 5 3. Apply pooled desalted serum to a pre-equilibrated (20ml application buffer) Synsorb 115 (galactosylgalactose; Chembiomed, Alberta, Canada) or D(+) Melibiose-Agarose (Sigma) affinity column (5ml-50 ml depending on the yield required).
- 4. Collect run-through (partially anti-GAL-depleted) and reapply to column. Repeat process 3 times to ensure complete removal of anti-GAL antibodies. The wash-through from the 3rd passage through the Synsorb column is collected and the volume adjusted to the original volume of the serum with phosphate-buffered saline (PBS) pH 7 +0.05% azide. This is used as a source of anti-GAL
 - antibody-depleted serum.

 5. Wash column with PBS pH 8 until the eluate is protein free (0.D. 280nm=0).
- 20 6. Elute anti-GAL antibodies with 3.5M KSCN, pH 7.5. Collect 4ml fractions, determine the O.D. 280 and pool peak fractions (usually 1-6).
- 7. Concentrate anti-GAL antibodies using CF25 ultrafiltration cones (Amicon, Danvers, USA). Add 7ml of 25 the pooled fractions containing anti-GAL antibodies to spin cone and centrifuge (2,000 RPM, 10min, 4°C). Refill cone and recentrifuge until volume is reduced to 3-5ml.
- 8. To dilute the KSCN, adjust vol. to 7ml with PBS and centrifuge (2,000 RPM, 10min, 4°C). Repeat process a 30 further 10 times.
 - 9. Remove sample containing anti-GAL antibodies from cone using plastic pipette; rinse cone with PBS pH7 +0.05% azide.

- B. <u>Preparation of IqG anti-GAL antibodies</u>
 The following procedures are performed at 4°C.
- Desalt 15-30 ml serum (in 3ml batches) by passage through a pre-equilibrated (20ml application buffer)
 Econo Pac 10DG (Bio-Rad, Richmond, USA) column.
 Alternatively for large scale preparations desalt by dialysis exhaustively against application buffer.
 - 2. Wash column with 4ml aliquots of application buffer. Collect and pool column eluates.
- 3. Apply desalted serum to a pre-equilibrated (30ml application buffer) Affi-Blue column (Bio-Rad, Richmond, USA) (Affi-Blue binds all proteins except albumin and IgG).
- Wash column with 20ml application buffer to elute
 15 IgG enriched fraction:
 - 5. Apply IgG enriched fraction to a pre-equilibrated (20ml application buffer, pH 8.0) Synsorb 115 (galactosyl-galactose; Chembiomed, Alberta, Canada) affinity column (5ml).
- 20 6. Collect run-through and reapply to column. Repeat process 3 times to ensure complete removal of anti-GAL antibodies. The wash-through from the 3rd passage through the Synsorb column is collected and the volume adjusted to the original volume of the serum with PBS pH 25 7 +0.05% azide. This is used as a source of control

anti-GAL-depleted IgG.

In some cases anti-GAL IgG was further purified using a protein G column, which efficiently binds IgG but not other antibody isotypes. IgG was then eluted from the protein G column using glycine pH 2.4.

All anti-GAL antibody preparations were analyzed for the following:

a. Protein content was determined using the Bradford colorimetric method (Bradford, M.M 1976,

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Anal. Biochem. 72:248-254), using purified human IgG as the standard.

- b. Molecular weight and purity were determined using polyacrylamide gel electrophoresis according to method described by Laemli, Nature (London) 227: 680 (1970), and protein was detected in the gels by silver staining using standard kit reagents (Amersham, UK).
- c. Antibody class and isotype were determined by radial
 immunodiffusion using standard techniques as set out in Rose et al. (eds.), Manual of Clinical Laboratory Immunology, American Society for Microbiology,
 Washington, D.C. IgG anti-GAL preparations were found to contain all subclasses, with IgG2 predominating.

25 EXAMPLE 2

Reactivity of IgG and IgM Anti-GAL Antibodies and
Depleted Serum with Porcine Cells and Tissues

I. CELLS

Reactivity of IgG and IgM anti-GAL antibodies was assessed using either porcine aortic endothelial cells (prepared by the inventors as described below) or porcine epithelial cell line LLC PK_1 (PK_1), obtained from the

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American Type Culture Collection (ATCC), Accession No. CRL1392.

A. <u>Isolation and culture of porcine aortic</u> endothelial cells (PAE's)

5 Pigs were blood typed (using human typing reagents) to identify "O-type" pigs, i.e, pigs unreactive with antibodies to A or B human red blood cell antigens. Aortas were excised from "0-type" pigs, then transported from the abattoir to the laboratory on ice. PAE's were 10 isolated by collagenase treatment as described by Gimbrone et al., J. Cell Biol. 60: 673-84 (1974). were cultured in RPMI medium containing 10% fetal calf serum (FCS), supplemented with $150\mu g/ml$ endothelial cell supplement (Sigma) and $50\mu g/ml$ heparin (Sigma). 15 cells were identified as endothelial cells by their typical cobblestone morphology and by their immunoreactivity with Factor VIII antibodies, as identified using immunofluorescence. In all the assays described below, the PAE's were used between the 8th and 20 12th passages.

B. Tissue Culture: Maintenance of PK-1 and PAE cell lines

All tissue culture was performed in a laminar flow hood, using appropriate tissue culture sterile technique.

25 All tissue culture reagents, unless otherwise indicated, were purchased from CSL, Melbourne, Australia. Media were constituted as follows:

PK-1 Culture Medium:

DMEM (Cytosystems, Castle Hill, Australia) 500ml
30 FCS (CSL, Melbourne, Australia) 37.5ml
Glutamine (200mM) (Cytosystems) 5ml
Hepes (1M) (CSL) 7.5ml
Penicillin (CSL) 0.5ml (10⁵U/ml final)
Streptomycin (CSL) 0.5ml (10⁵µg/ml final)

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PAE - Culture Medium:	
RPMI (CSL)	90ml
FCS (CSL)	10ml
Endothelial cell	
<pre>supplement (3mg/ml) (Sigma)</pre>	1.5ml
Heparin (10mg/ml) (CSL)	0.5ml

Endothelial cell supplement was purchased from Sigma Chem. Co. (St. Louis, Missouri, USA) as a lyophilized powder, resuspended in sterile HBBS, and 3ml aliquots stored at 4°C.

Heparin (Sigma, Missouri, USA) - dissolved in PBS (10mg/ml) - filter sterilized (0.22vm)

15 Hanks Buffer - purchased from Cytosystems

The following general procedures were used in propagating the cell lines.

Pour off old medium
 Rinse cells twice with sterile PBS
 Add 3ml of TED (0.05 M trypsin, 0.53 M EDTA, Gibco, NY,USA)
 Incubate 10 min. in CO₂ incubator at 37°C

5) Add 7ml RPMI with 10% FCS

- 25 6) Resuspend cells and transfer to a sterile 10ml tube
 - 7) Centrifuge for 5min at 1200 rpm, discard supernatant
 - 8) Resuspend cells in RPMI with 10% Newborn Bovine Serum (NBS) and repeat centrifugation
 - 9) Resuspend cells in 1ml DMEM (PK-1's) or RPMI (PAE's) (with additives, as described above).
- 10) Add 10ml medium and the appropriate volume of cell suspension to achieve the desired dilution for each 75cm² tissue culture

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flask, and return to humidified CO_2 incubator.

- C. Antibody staining and FACS analysis
- 1) Add 2ml TED to a 75cm² culture flask containing PK-1 or PAE's, and incubate at room temperature for 10 min.
- 2) Add RPMI plus 10% FCS (5ml) to neutralize trypsin.
- Pellet cells by centrifugation (700g, 5 min, 4°C).
 - 4) Wash cells by resuspension and centrifugation in Hanks Buffer (x2).
 - 5) Pellet cells by centrifugation (700g, 5 min, 4°C).
 - Resuspend cell pellet in Hanks buffer containing purified anti-GAL antibodies, GAL-depleted serum or GAL-depleted IgG and incubate at 4°C for 60 min. All antibodies were used undiluted, or diluted 1:2 or 1:4 in Hanks buffer.
 - 7) Add 1ml of Hanks Buffer, pellet cells by centrifugation and aspirate off supernatant.
 - 8) Resuspend pellet in FITC-labelled sheep-antihuman IgG Fab2 or IgM Fab2 (Silenus, Hawthorn, Australia) diluted 1:80 in Hanks buffer.
 - 9) Incubate for 30 min. at 4°C.
 - 10) Wash three times with Hanks buffer; resuspend pellet from final wash in 0.5ml Hanks buffer.

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11) Analyze stained samples using a FACScan II (Becton Dickinson) according to the manufacturer's instructions.

The specificity of the anti-GAL antibody binding 5 to porcine cells was determined by examining the ability of sugars of various structures to inhibit antibody In these competition studies the anti-GAL antibodies were pre-incubated with sugar (0.1M) at 37°C for 30 min before adding to the cells.

D. Results

10 Using immunofluorescence it was found that total anti-GAL (IgM & IgG) and purified anti-GAL IgG stained both PK-1 and PAE's cells. On the other hand, the total anti-GAL antibody-depleted serum nor the anti-15 GAL IgG-depleted serum gave detectable staining over background. The staining with anti-GAL IgM and/or IgG was inhibited with purified galactose and with disaccharides having terminal galactose residues in the α 1-configuration such as melibiose (6-0- α -D-20 galactopyranosyl- D-glucose) and stachyose (α -D-Gal-[1->6]- α -D-Glc-[1->2]- β -D-Fru). Staining was not inhibited with sugars such as lactose $(4-0-\beta-D)$ -galatopyranosyl- $\alpha-D$ glucose), which has a terminal galactose residue, but in a β 1->4 configuration. The results of one such 25 experiment are represented in Figure 1. PAE's were stained with anti-GAL antibody alone (GAL:PBS) or with anti-GAL antibody that had been pre-incubated with either melibiose (GAL:MELIBIOSE), galactose (GAL:GALACTOSE) or

lactose (GAL:LACTOSE). Anti-GAL antibody staining was 30 approximately 10 fold less in the samples containing melibiose and galactose, but was not affected significantly by lactose.

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II. TISSUES

A. Methods

Pig kidney was fixed in formalin and dehydrated before embedding in Paraplast. Pig heart and liver were 5 fixed in paraformaldehyde-lysine-periodate fixative and snap frozen in O.C.T. embedding compound (10.24% w/w polyvinyl alcohol, 4.26% w/w polyethylene glycol, 85.50% w/w nonreactive ingredients; Tissue Tek®, Miles, Inc., Elkhart, Indiana, USA). Four μ m-thick sections of pig 10 heart and liver and 2 μm -thick sections of kidney were incubated with purified anti-GAL antibodies (undiluted, 1:2 and 1:4) for 60 min. and then incubated with a fluorescein isothiocyanate (FITC)-conjugated sheep antihuman immunoglobulin F(ab') fragment (Silenus 15 Laboratories, Hawthorn, Australia) (1:100) for 30 min. or a peroxidase-conjugated rabbit anti-human IgG (Dakopatts, Glostrup, Denmark) (1:50) for 60 min. Control sections were analyzed for autofluorescence, with the secondary antibody alone, or with the anti-GAL-depleted IgG or 20 normal pig serum as the primary antibody. No staining was detected. The specificity of the anti-GAL antibodies was tested by pre-incubating sections of pig renal cortex with a variety of sugars, including melibiose, lactose, sucrose and glucose at 0.1M.

B. Results

As with the analyses performed on the pig cells using immunofluorescence, total anti-GAL IgM + IgG, purified anti-GAL IgG, but not the anti-GAL IgM and/or IgG-depleted sera, stained all pig tissues examined. The individual staining parameters varied from organ to organ as set out below:

- 41 -

Immunostaining of Pig Tissues with Anti-GAL Antibodies:

	<u>Tissue</u>	Anti-GAL Reactivity	<u>Stain</u>	ing Intensity
5	Kidney	Proximal and distal convoluted tubules Endothelium: Intertubular sinusoids Endothelium: Arteries and veins Glomerular capillaries		Variable Variable Strong Variable
	Heart	Endothelium: Arteries, veins, capillaries Endocardium Myocardium	3 .	Strong Strong Perinuclear
10	Liver	Small Bile Ducts (lining cells) Endothelium: Arteries, veins Intertubular sinusoids		Strong Strong Negative

The specificity of the binding of anti-GAL antibodies was tested on sections of pig renal cortex by inhibition with 0.1 M melibiose, lactose, sucrose and glucose. Reactivity of the anti-GAL antibodies with proximal tubule brush borders was reduced to near background by preincubation of antibody with melibiose, but was not inhibited by the other saccharides.

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EXAMPLE 3

Hemagglutination of Pig RBC by Human Serum: Sugar Inhibition Studies

The methods used to investigate the hemagglutination of pig red blood cells (RBC's) by human 25 serum was adapted from the methods described by Galili, J.Exp. Med. 160: 1579-81 (1984) and Severson, Immunol. 96: 785-789 (1966).

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I. METHODS

A. <u>Media/Solution Preparation</u>

- Human Serum Albumin (HSA) (CSL, Melbourne, Australia) (5mg/ml) was dissolved in PBS,
 filter sterilized, and stored at 4°C.
 - 2. Preparation of sugars:
 1M stock solutions of sugar were
 prepared by dissolving the amount
 indicated in 100ml of PBS. Sodium azide
 was added (0.02%) and solutions stored at
 4°C.

	α -Lactose (4-O- β -D-galactopyranosyl- α -D-glucose D(+)galactose	36.0g 18.0g
	Stachyose $(\alpha-D-gal-[1->6]-\alpha-D-Glc-[1->2]-\beta-D-Fru)$	66.6g
15	Melibiose (6-O- α -D-galactopyranosyl- D-glucose) Sucrose (α -D-Glucopyranosyl β -D-fructofuranoside) D-(+)-Glucose α -D-(+)-Fucose (6-Deoxy-D-galactopyranose)	34.2 g 34.2 g 18.0 g 16.4 g

All sugars were purchased from Sigma (St. Louis, 20 Missouri, USA). Sugar solutions were diluted in PBS to the appropriate concentration as required.

B. <u>Preparation of pig RBC's</u>

- 1. Heparinised pig blood
 (Animal Resources, Clayton,
 Australia) is centrifuged at 800
 RPM for 10min to pellet the RBC.
- 2. The RBC pellet is washed by resuspension in PBS (10ml) and recentrifugation (repeated 3 times). After the final wash, the RBC pellet is resuspended in 10ml PBS.
- 3. A 0.5% solution of RBC's is prepared by adding 50ul RBC solution (from step 2, above) to

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10 ml PBS containing 0.5g/100 ml of HSA.

- C. <u>Preparation of 96-well microtitre plates</u>
 (Titretek, USA)
- Add 25ul of PBS to each well.
 - 2. Add 25ul of pooled human AB serum (CSL, Melbourne, Australia) to column 1 and serially dilute by removing 25ul from column 1 and adding to column 2, then repeating by sequentially removing and adding 25ul from and to each well across the plate, finally discarding 25ul from column 11 and adding no serum to column
- 3. Add 25ul of sugar solution to each row in decreasing concentrations down rows. No sugar solution is added to the final row.
- 4. Incubate at 4°C overnight
 25 and then at 37°C for 30 min.
 5. Add 50ul of 0.5% pig RBC to
 each well; vortex and incubate
 at room temperature for 2 hours.
 Determine agglutination
- 30 visually.

12.

II. RESULTS

Human serum caused the agglutination of pig RBC's at a titre of between 1/32-1/64, which is consistent with the presence of high levels of naturally occurring

- 44 -

xenoantibody (NXAb) in human serum. To examine the specificity of the NXAb response, sugar inhibition studies were performed. Sugars such as melibiose, stachyose, galactose and fucose which have terminal galactose residues in the α1-6 configuration were found to inhibit agglutination in the μM to mM range. Sugars with other structures, such as lactose and sucrose, were only inhibitory when very high concentrations were used. At these high concentrations, the observed effects are most probably non-specific, due, for example, to changes in osmolarity. Results are summarized below:

Pig RBC Hemagglutination by Human Serum: Sugar Inhibition

	Sugar	Linkage	Inhibitory Concentration
	Melibiose	Gal α 1-6Glc	5x10 ⁻⁴ M
15	Stachyose	Gal α 1-6Gal	2x10 ⁻³ M
	Galactose		2×10 ⁻³ M
	Fucose	6-Deoxy-α-L-Gal	1x10 ⁻³ M
	Lactose	$Gal\beta1-4-Glc$	> 10 ⁻¹ M
	Sucrose	α-D-Glc-β-D-Fruc	>10 ⁻¹ M

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EXAMPLE 4

Inhibition of Human Serum-Induced Lysis of Porcine <u>Cells by Sugars</u>

The ability of human serum to cause the lysis of porcine cells was examined using both pig epithelial

25 (PK₁) and aortic endothelial (PAE's) cells, the isolation and culture of which is described in Example 2. Cell lysis was determined using either the ⁵¹Chromium release assay as described by Cerottini and Brunner, Nature New Biol. 237:272, 1972 or the Cytotox LDH release assay according to the manufacturer's instructions (Promega, USA).

I. METHODS

- A. 51 CR Release Assay
- Preparation of Cells:

- 45 -

- a) Trypsinize a confluent flask of cells. On average, approximately 3 x 10^6 PAE's and approximately 3 x 10^7 PK₁ cells are obtained per 10 ml flask. About 1 x 10^5 cells are required for each well in the 51 CR Release 5 Assay.
 - b) Wash cells 4 times in 10 ml RPMI (no FCS); spin 1200 rpm for 5 min.
 - c) Resuspend cells in 100 μ l RPMI (with 10% heat-inactivated FCS; see below).
- 10 2. Labelling Cells with 51 CR:
 - a) Combine in a 10 ml tube: Cells in 195 μ l RPMI/10% FCS (heat inactivated); 5 μ l 51 CR (120 μ Ci).
 - b) Incubate at 37°C for 2 hr.
 - c) Add 2 ml RPMI/10% FCS (heat
- 15 inactivated).

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- d) Centrifuge cells through a layer of FCS (heat inactivated) to remove excess label.
- e) Gently overlay the labelled cells onto a 4 ml cushion of FCS using a Pasteur pipette.
 - f) Centrifuge at 700g for 5 min. at 4°C.
- g) Remove supernatant taking care not to disturb the cell pellet.
- h) Resuspend pellet in RPMI/10% FCS (heat inactivated) at about 3 \times 10⁷ cells/ml.
- 25 3. Assay Conditions:
 - a) For PAE's, rabbit complement was used as the complement source, since the ⁵¹ CR-release assay was not sufficiently sensitive to detect lysis when human complement, a less "active" source, was used. In
- 30 contrast, with the LDH assay, which is significantly more sensitive, normal human serum (NHS) was used as the source of complement.
 - b) To each test well of a 96-well V bottom plate, add:
- 100 μ l labelled cells

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- 10-50 μ l NHS (heat inactivated) (5-25%

of final)

- Complement:

PAE's: 50 μ l absorbed

5 rabbit complement (25% final)

 PK_1 : 10-40 μ 1 NHS (5-25% of

final)

- 50 μ l antibody (total anti-GAL (IgG + IgM, anti-GAL IgG, anti-GAL antibody-depleted serum, or 10 anti-GAL antibody-depleted IgG)
 - Adjust volume to 200 μ l with C) RPMI/10% FCS (heat inactivated) if required
 - 'd)' Incubate plates at 37°C for 3 hr.
 - Centrifuge plates at 1000 rpm for 5
- 15 min to pellet cells
 - f) Remove 100 μ l of supernatant from each well and transfer to a gamma counter tube
- Add 3 ml scintillation fluid and g) measure ⁵¹ CR release using a gamma counter (Packard 20 Instrument Company, Illinois, USA)

(To determine maximum release, add 100 μ l 8% Triton X-100 made up in RPMI/10% FCS (heat inactivated) to 100 μ l labelled cells)

(Note: Each reaction is set up in

- 25 quadruplicate)
 - Calculation of % Lysis:
 - % Lysis = Experimental cpm Spontaneous Release cpm x 100 Max. Release cpm - Spontaneous Release cpm
- 5. Sugar Inhibition of Complement-Induced Cell 30 Cytotoxicity:

In a 96-well test plate, mix the following:

- 50 μ l labelled cells

- 50 μ l complement

(PAE's: pig spleen cell absorbed complement; PK1's: NHS)

- 35 - x μ l sugar (final concentration of sugar: 10^{-1} to 10^{-3} M)
 - y μ l NHS (heat inactivated) final concentration 5-20%)
 - make volume to 200 μ l with RPMI

Plate Layout:

 Plate 1
 Plate 2

 5% 10%
 15% 20%

Rows:

1-4 5-8

1-4 5-8

Columns:

- 1. Spontaneous Release
- 2. Maximum Release
- 3. Melibiose
- 4. Lactose

B. <u>LDH Release Assay</u>

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- 1. General Procedures:
- a) Prepare cells as for ⁵¹ CR Release assay, and labeled with LDH as per the manufacturer's instructions (Cytotox non-radioactive LDH release assay, Promega, USA)
- b) To each well of a 96-well plate add (each reaction set up in quadruplicate):
 - 25 μ l labeled cells
 - 5-20 μ 1 NHS
 - $x \mu l$ sugar (final concentration of sugar: 10^{-1} to 10^{-3} M) RPMI/10% FCS (heat inactivated), to total
 - volume of 100 μ l
 - c) Incubate plates at 37°C for 3 hr.
 - d) Centrifuge plates at 1500 rpm for 5 min.
- e) Remove 50 μ l supernatant from each well (taking care not to remove any cells) and transfer to ELISA plate containing 50 μ l substrate mix (prepared according to manufacturer's instructions
- f) Cover tray and incubate in the dark 30 at room temperature for 30 min.
 - g) Add 50 μl stop solution to each well using multichannel pipette
 - h) Read absorbance at 492 nm.

2. Controls:

PCT/IB95/00088

- a) Spontaneous release (no antibody or complement)
 - 25 μ l labeled cells
 - 75 μ l RPMI/10% FCS (heat inactivated)
- 5 b) Maximum release
 - 25 μ l labeled cells
 - 50 μl 16% Triton X-100
 - 25 µl RPMI/10% FCS (heat inactivated)
 - Calculation of % Lysis: % Lysis =
- 10 Experimental release (Spontaneous release cpm + sugar cpm) x 100 Maximum release - (Spontaneous release cpm + sugar cpm)
 - Experimental Design:

```
Plate 1
```

15 Columns: 1.spontaneous release Rows: 1-4: cells + no sugar 5-8: no cells + no sugar

maximum release

3. 5% serum

4. 10% serum 5. 25% serum

20 6. RF10 alone

> Plate 2 melibiose Plate 3 galactose Plate 4 lactose

Plate 5 sucrose 25 Plates 6-9 same as plates 2-5 but no cells added

Sugar Conc. 1 x 10⁻¹M 5 x 10⁻²M Columns: 1-2

Rows: 1-2 O% serum 3-4 3-4 5% serum 5-6 1 x 10-2M 7-8 5 x 10-3M 9-10 2 x 10-3M 10% serum 5-6 30 7-8 25% serum

11-12 1 x 10⁻³M

- 5. Preparation of Pig Spleen-Absorbed Rabbit Complement:
- a) Cut pig spleen (obtained from local abattoir) into small pieces and prepare a single-cell suspension by passage through a fine metal sieve
- b) Pellet cells by centrifugation at 700g, 7 min. at 4°C
 - c) Resuspend cell pellet in RPMI/10% FCS and repeat centrifugation
 - d) Resuspend in RPMI/10% FCS/10%
- 10 dimethylsulfoxide (DMSO)
 - e) Count cells and store frozen aliquots (3 \times 10 9 cells/aliquot)
 - use one aliquot for each absorption
- f) For absorption, thaw and centrifuge at 600g, 5 min. at 4°C and remove the supernatant containing the DMSO
 - g) Wash two times with RPMI/10% FCS (10 ml)
 - h) Resuspend the cell pellet in rabbit complement; mix (rotary wheel) 2 hr. at 4°C
- i) Centrifuge 600g, 5 min. at 4°C and remove the supernatant containing the rabbit complement; store at 4°C

II. RESULTS

Comparable results were obtained with both cell types (PAE's and PK₁'s) using both lysis assays. The results of a typical lysis experiment are represented in Figure 2, in which the lysis of PAE's by human serum and by purified anti-GAL antibodies was determined using the ⁵¹CR release assay. Comparable results were also obtained with PK₁ cells using the ⁵¹CR release assay and with both cell lines using the LDH release assay. The results of

1. Xenoantibodies (NXAb) in human serum in the presence of complement are capable of lysing porcine

these assays can be summarized as follows:

cells. Lysis increases with increasing concentrations of serum.

- 2. Pre-absorption of NHS with pig spleen cells (which removes the NXAb): <u>No lysis</u>.
- 5 3. Use of heat-inactivated complement: No lysis.
 - 4. Use of NHS depleted of anti-GAL antibodies: No lysis.
- 5. Use of purified total anti-GAL antibodies 10 (IgG + IgM): <u>Lysis</u>.
 - 6. Use of purified anti-GAL IgG: No lysis.
- Use of purified total anti-GAL antibodies
 (IgG + IgM) and dithiothreitol (DTT): No lysis. (DTT is
 a reducing agent that disrupts the multimeric structure
 of IgM antibodies without affecting IgG.)

Together these results demonstrate that the anti-GAL antibodies are responsible for the observed lysis. Purified anti-GAL IgG and DTT-treated total (IgG + IgM) anti-GAL antibodies failed to elicit lysis, indicating that IgM, but not IgG, antibodies are causative agents in this system. Preliminary attempts to verify this observation using purified IgM prepared either in crude form by euglobulin fractionation or by α-IgM affinity chromatography were unsuccessful. The inventors believe this reflects inactivation of the IgM during preparation, rather than a true reflection of the capacity of anti-GAL IgM to cause lysis of porcine cells. heat inactivation of the complement prevented lysis, indicating that lysis of porcine cells is a complement-dependent phenomenon.

The effect of adding the disaccharide sugars melibiose (Gal α 1 \rightarrow 6 Gal) and lactose (Gal β 1 \rightarrow 4 Glu) on the lysis of PAE's by human serum was assessed using the Cytotox non-radioactive LDH release assay. PAE's were incubated in the presence of 50% human serum as the

source of xenoantibody and complement, together with various concentrations of each sugar (1mM to 100mM). Under these conditions, melibiose, which has the Gal α1→6 Gal configuration, but not lactose, which has the terminal Gal moiety by in a β1→4 configuration, protected the pig cells from lysis.

EXAMPLE 5

Inhibition of Human Serum-Induced Damage to Rat Hearts by Sugars

The Langendorf isolated perfused ex vivo heart

model was used to further demonstrate the involvement of anti-GAL xenoantibodies in hyperacute rejection.

I. METHODS

- A. Preparation and storage of Human Plasma
 - 1. Centrifuge fresh human blood at 3000
- 15 rpm, 10 min., 4°C to remove red blood cells (RBC's)
 - 2. Remove the plasma
 - 3. Centrifuge the plasma at 10,000 rpm, 10 min. 4°C to remove any remaining cells; decant the plasma
 - 4. Add 2.5 ml of 0.1M EDTA pH 7.30 for
- 20 every 50 ml of plasma
 - 5. Store 50 ml aliquots at -70°C
 - 6. For heat-inactivated plasma, heat at 56°C for 60 min., then centrifuge at 2,500 rpm for 10 min.
- B. <u>Assessment of Complement Activity</u>

Before being used in the <u>ex vivo</u> model, both heat inactivated and control plasma was tested for complement activity. Classical complement activity was determined by hemolysis using sensitized sheep RBC's as described by

30 Harrison and Lachman, In: Weir et al. (eds.), <u>Handbook</u>
of Experimental Immunology and Immunochemistry, 4th Ed.,
Blackwell scientific Publications (1986). Alternative
complement pathway activity was determined using the

- 52 -

rabbit hemolytic assay as described by Serrais et al., J. Immunol. Meth. 140: 93-100 (1991). The assay was performed in buffer containing EGTA and MgCl₂. The EGTA chelates the Ca⁺⁺, thus inhibiting the classical pathway.

5 The Mg⁺⁺ is required for activation and assembly of CdbBb, the alternative pathway C3 convertase.

- C. Preparation of Plasma for Heart Perfusions
 Plasma prepared from different blood packs is
 thawed at 37°C, pooled and filtered (100 μm steel mesh,
 8.0 μm and 4.5 μm Millipore filters, sequentially).
 CaCl₂ is added at 0.58 mg/ml plasma, and the plasma kept
 on ice until ready for perfusion.
 - D. <u>Ex Vivo Isolated Perfused Rodent Heart Model</u>
- 1. Anesthetize rats with Nembutal (1 μ l sodium 15 pentobarbitone (60 mg/ml)/g body weight) and mice with ether.
 - 2. Surgically expose the heart and inject heparin (Porcine Mucous, 10,000 U/ml) into the femoral vein (rats: 0.3 ml injected).
- 3. Remove heart and place in ice-cold Krebs-Henseleit buffer containing heparin (0.2 ml/50 ml buffer. Krebs-Henseleit buffer:

- 119 mM NaCl - 25 mM NaHCO₃ - 4.6 mM KCl - 1.2 mM MgSO₄·7H₂O - 1.3 mM CaCl₂·2H₂O - 1.2 mM KH₂PO₄ _ 11 mM glucose - 0.25% (v/v) BSA

- Adjust to pH 7.4; store at 4°C

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4. Connect aorta to the canula of the Langendorf perfusion apparatus and tie firmly. The apparatus was

assembled by the present inventors according to

35 experimental requirements of the Langendorf heart model
as described in Doring & Dehnerrt, The Isolated Perfused
Heart According to Langendorf, Bionesstechnik-Verlag
March GmbH, D7806, West Germany.

- 5. Perfuse with Krebs-Henseleit buffer (made fresh each day), which is gassed continuously with carbogen (95% O_2 , 5% CO_2) at a pressure of 100 mmHg, at 37°C.
- 6. Attach a hook, connected to a transducer (Physiograph MK-111-S, Narco Bio-Systems) to the apex of the heart.
- 7. Perfuse heart for 20 min. with Krebs-Henseleit buffer to enable heart to stabilize (reservoir 10 volume: 270 ml).
 - 8. Add plasma (pre-warmed to 37°C) as follows:
 - at 20 min. add 10 ml plasma (= 5% plasma)
 - at 25 min. add 10 ml plasma (= 9 % plasma)
 - at 30 min. add 10 ml plasma (= 13 % plasma)
- 9. Monitor heart for a further 30 min. and record heart flow and contraction rate.

E. Sugar perfusion

- 1. Stabilize heart in Krebs Henseleit buffer for 30 min. as described above.
- 2. Add 2.5 ml of 1.08 M stock sugar solution to reservoir; total volume = 270 ml; final sugar concentration = 10mM.
- 3. Allow heart to restabilize for 10 min, then add plasma (control or heat inactivated) as per the 25 schedule described above.
 - 4. Record heart beat and flow rate.
 - F. <u>Large-Scale Preparation of anti-GAL antibody-</u>
 <u>Depleted Plasma</u>

(all manipulations are performed at 4°C)

1. Start with 200 ml freshly prepared human plasma; 100 ml is subject to depletion; 100 ml is used as an untreated control from the same patient drawn on the same day; store at 4°C.

- 2. Filter the plasma sequentially through a 100 μm , 8 μm metal sieves and finally through a 0.45 μm Millipore filter; dilute to 1000 ml with PBS, pH 8.0.
- Concentrate to 200 ml using an Amicon spiral
 wound cartridge (removes salt).
 - 4. Equilibrate melibiose sepharose column (40 ml) with PBS, pH 8.0 (10 column volumes).
- 5. Passage the plasma through the melibiose sepharose column; collect the run-through and store at 10 70°C (=partially depleted plasma).
 - 6. Wash column with PBS, pH 8.0 (10 column volumes) until the O.D. (280nm) of the eluate is approximately zero.
- 7. Combine the partially depleted plasma and the 15 eluate from the wash; concentrate to 200 ml (Amicon spiral concentrator).
 - 8. Elute the anti-GAL antibody fraction with 4M guanidinium HCl pH 6.4 (2 column volumes).
- 9. Regenerate the column with PBS (10 column 20 volumes).
 - 10. Repeat the entire process an additional two times, i.e., repassage plasma through the melibiose column, wash, elute the anti-GAL antibody fraction and regenerate column.
- 25
 11. For the anti-GAL antibody-depleted fraction:
 combine the eluate from the melibiose sepharose column with run-through from the final wash
 adjust the volume to 5 liters with Krebs Henseleit buffer and add EDTA to 10 mM; adjust pH
 30 to 7.0
 - concentrate back to original volume (Amicon spiral concentrator); aliquot (35 ml) and store at -70°C
 - 12. For the anti-Gal antibody fraction:

- combine the eluted anti-GAL antibody fractions, dilute to 5 liters with Krebs Henseleit buffer and add EDTA to 10 mM
- concentrate back to 10 ml (Amicon 5 spiral concentrator); aliquot (1 ml) and store at -70°C 13. The anti-GAL antibody-depleted fraction and the purified anti-GAl antibody fraction are tested for
- a) Anti-GAL reactivity: Use as primary reagents to stain porcine cells (PK1's). Detect 10 staining as described in Example 2, above. Analyze stained samples using a FACScan II (Becton Dickinson), according to the manufacturer's instructions.
- b) Protein content: Determine using the colorimetric method of Bradford, Anal. Biochem. 72:
 248-54 (1976), with purified human IgG as the standard.
- c) Electrolyte concentration: On the day of the perfusion, the anti-GAL antibody depleted plasma is also tested to determine the calcium, magnesium and potassium levels using an electrolyte autoanalyser (Olympus); the levels of each are adjusted to normal as required.

II. RESULTS

Rat hearts were connected to the Langendorf apparatus and then stabilized by perfusion with Krebs

25 Henseleit buffer for 10 min., and then a further 10 min. with the same buffer containing either melibiose or lactose (10mM). Human plasma was then added in stages as described above to a final concentration of 13 % and the effect of the added sugar on cardiac function was

30 assessed. The parameters measured were heart rate, amplitude (strength) of contraction and output (Figure 3).

In the presence of human serum alone (lower trace), the heart essentially stopped beating within

minutes. The same result was obtained if lactose was added. In the presence of melibiose (upper trace) or anti-GAL antibody-depleted plasma, however, the heart was able to maintain a strong beat. When the purified anti-GAL antibody was added back to the anti-GAL antibody-depleted plasma, the heart again stopped beating within minutes.

EXAMPLE 6

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Characterization of the Porcine α -1,3-GalT Gene cDNA's encoding porcine α -1,3-GalT were generated 10 by Polymerase Chain Reaction (PCR) technology. Total RNA of pig liver was isolated by homogenizing liver slices in 7M guanidinium thiocyanate, as described by Chomczynski & Sacchi, Anal. Biochem 162, 156-159 (1987); Sambrook et 15 al., Molecular Cloning: A Laboratory Manual (2nd Edition), Cold Spring Harbor Laboratory Press (1989). Sixteen μ g of the RNA, together with 1μ g oligo dT primer, were heat denatured for 5 minutes at 65°C prior to being transcribed into cDNA using avian myeloblastosis virus 20 (AMV) reverse transcriptase in a 100 μ l reaction carried out at 37°C for 90 minutes. Three μ l of the cDNA synthesis reaction was used in the subsequent PCR amplifications. General procedures used for generation of cDNA are provided in Sambrook et al (1989), supra.

Primers for PCR were synthesized using phosphoramidite technology, on an Applied Biosystems DNA synthesizer. The sequence of the PCR primers was based on identifying conserved regions within the published sequences for murine and bovine $\alpha-1,3$ -GalT genes.

Joziazze et al., J. Biol. Chem 264: 14290-97 (1989);
Joziazze et al., Biol. Chem 267: 5534-5541 (1992). All
primers were synthesized with EcoR1 linkers at the 5' end
for ease of cloning. In the following listing of the
primers used in the present study, nucleotide positions

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varying between bovine and murine sequences are singleunderlined; nucleotide positions varying between bovine and human sequences are double-underlined:

Exon 2 primer (forward):

5 5'-GTGAATTCAGCCCTGCCTCCTTCTGCAG-3'

(SEQ ID NO: 1)

Designation:

GTE2F -- 28-mer

- 1 difference b/w bovine & murine
- no sequence available for human exon 2
- 10 Exon 4 primer (forward): 5'-GTGAATTCAGGAGAAAATAATGAATGTC-3'

(SEQ ID NO: 2)

Designation:

GTE4F -- 28-mer

- no differences b/w bovine, murine & human
- 15 Exon 9 primer (reverse): 5'-GTGAATTCGGGA<u>TCT</u>GCCTTGTACC<u>A</u>CC-3'

(SEQ ID NO: 3)

Designation:

GTE9R -- 28-mer

- 3 differences b/w bovine & murine
- 20 1 difference b/w bovine & human

3'-UTR primer (reverse):
5'-GTGAATTCGAAATCACTGGGAATTTACA-3'

(SEQ ID NO: 4)

Designation:

GT3UR -- 28-mer

- no differences b/w bovine & murine

- no differences b/w bovine & human

Exon 9 primer (forward): 5'-AGGAATTCAGCATGATGCGCATGAAGAC-3'

(SEQ ID NO: 5)

30 Designation:

GTE9F -- 28-mer

- no differences b/w bovine & murine
- 3 differences b/w bovine & human

- 58 -

PolyA primer (reverse): 5'-TTGAATTCTTTTTTTTTTTTTTTTTV*N**-3'

(SEQ ID NO: 6) * V = A or C or G; ** N = A or C or G or T

(primer includes all nucleotide variants
5 for V and N)

Designation: APATR -- 23-mer

The PCR conditions used to generate porcine α -1,3-GalT cDNA fragments were as

follows:

MICHACID- -MIC

- 10 1) For GTE2F + GTE9R and GTE4F + GTE9R: heat to 94°C (60 seconds); then proceed with 35 reiterations (cycles) of the following three steps: (1) 94°C, 40 seconds, (2) 57°C, 50 seconds, and (3) 72°C, 80 seconds.
- 2) For GTE9F + GT3UR: heat to 94°C (120 seconds); then
 15 proceed with 35 cycles of: (1) 94°C, 40 seconds, (2)
 48°C, 45 seconds, and (3) 72°C, 60 seconds.

The PCR fragments were subcloned into EcoR1restricted pBluescript II KS+ (Stratagene, Cat, # 2
12206) and the DNA sequence was determined using the
chain termination method. The DNA sequence was assembled
and analyzed using DNASIS-Mac v2.01 (Hitachi)

The nucleotide sequence of porcine α-1,3-GalT (SEQ ID NO: 7) and the derived amino acid sequence (SEQ ID NO: 10) of the enzyme are shown in Figures 4 and 5. A single large open reading frame extends from the initiating methionine at nucleotide 91 to a stop codon located at nucleotide 1204. The sequence surrounding the putative initiating methionine conforms to the consensus eukaryotic initiation sequence. Kozak, Cell 44, 283-92 (1986).

The porcine cDNA sequence is compared to the corresponding murine (SEQ ID NO: 9) and bovine (SEQ ID NO: 8) sequences in Figure 4. The locations of introns within the murine gene are also shown. Joziazze et al., 35 J. Biol. Chem 267: 5534 (1992). This alignment

demonstrates that exon 3, located within the 5' untranslated region of the mouse gene, is not found in either the porcine or bovine cDNAs. The overall sequence identities between the coding sequences are as follows:

- 5 a) pig compared to mouse: 75.02% (exon 3 not considered)
- b) pig compared to bovine:- 85.15%
 The amino acid sequences of the porcine (SEQ ID NO: 10), murine (SEQ ID NO: 12) and bovine (SEQ ID NO: 11) α-1,3-GalT enzymes are depicted in Figure 5. The locations of introns are also shown, based on their positions within the mouse gene (Joziasse et al., 1992). This alignment illustrates that the overall amino acid
- a) pig compared to mouse: 71.98%

homologies are:

- b) pig compared to bovine: 82.87%
- c) bovine compared to mouse: 73.72%

EXAMPLE 7

Identification of Potential Sites to Interrupt the α -1-3-GalT Gene 20 The present inventors' choice of a site for interrupting the $\alpha-1,3$ -GalT gene has been influenced by several characteristics of the gene and its expression. In particular, several mRNAs for α -1,3-GalT have been detected in the mouse. Joziazze et al., J. Biol. Chem. 25 267: 5534 (1992). These mRNAs are products of alternative splicing events in which exons 5 and/or 6 may be deleted. Hence, these exons are not appropriate interruption sites in the mouse, since a transcript encoding a functional α -1,3-GalT enzyme presumably could 30 be formed when exons 5 or 6 are spliced out. Moreover, the present inventors have isolated two different classes of α -1,3-GalT cDNA clones from the pig - one that includes exon 5 and one with exon 5 deleted. It is possible that mRNA's with and without exon 6 are also

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formed by alternative splicing in the pig. Thus, for initial experiments the present inventors have not chosen these exons as sites for interruption.

Insertion of an interrupting-DNA fragment into

5 exon 4 (which encodes the cytoplasmic NH₂-terminal domain
and the membrane-anchoring domain; see Figure 5) would
disturb production of a transcript encoding an active α1,3-GalT. Hence this exon is an appropriate site to
disrupt the α-1,3-GalT gene. Similarly, exons 7 and 8,

10 which encode the NH₂-terminal region of the catalytic
domain, are suitable disruption sites. Insertion of a
interrupting DNA fragment within these exons would
prevent the synthesis of an active catalytic domain.

A preferred site for interrupting the mouse gene
15 is located at a Sall site found within exon 9 of the
mouse α-1,3-GalT gene, at codons 221 + 222 (see Figure
5). This site is positioned 150 amino acids from the
COOH-terminus, within the catalytic domain. The mouse
gene within the present inventors' constructs for
20 homologous recombination is interrupted at this Sall
site. The amino acids encoded by nucleotides at this
Sall site are conserved in the pig and bovine sequences,
although the Sall site itself is not. Construction of a
Sall site at this position in the pig gene (e.g., by in
25 vitro mutagenesis) provides a useful construct to
inactivate the gene.

EXAMPLE 8

Choice of a DNA Fragment to Interrupt the α -1,3-GalT Gene The present inventors have used both the neomycin

resistance (neo^R) gene and the hygromycin resistance gene (hyg^R) to interrupt the α-1,3-GalT gene. In one set of "knockout" constructs the neo^R and hyg^R genes are linked to the murine phosphoglycerate kinase (PGK) promoter (Adra et al., Gene 60: 65-74 (1987) and are both bordered by polylinker sequences that include restriction sites for EcoRV and ClaI.

In another construct, expression of the neo^R gene is directed by an altered polyoma virus promoter (PMC1; 10 Thomas and Cappechi, cell 51: 503-12 (1987)). In this construct the present inventors have addressed the problem of including an antibiotic resistance gene within the genome of transplant organs. That is, in some circumstances it may not be desirable to have genes conferring resistance to antibiotics present in the organ to be transplanted. The FLP/FRT recombinase system of yeast has been used to eliminate the neo^R gene from the sequence that interrupts the $\alpha-1,3$ -GalT gene.

In a construct of the present invention, the neo^R

20 gene is bordered at both the 5' and 3' ends by FRT DNA
elements. In addition, stop codons for each of three
reading frames have been inserted 3' to the neo^R gene,
and these stop codons, together with a single FRT
sequence, will remain within the α-1,3-GalT gene after

25 the neo^R gene has been excised by FLP. Targeted cells
carrying a genomic copy of the neo gene flanked by direct
repeats of the FRT could be supplied with FLP recombinase
in two ways:

1) Introduction into cells of partially purified 30 FLP protein:

FLP protein (0.1 - 10 μ g) is introduced ("transfected") into approximately 10⁷ cells using standard electroporation conditions. The cells are plated out into gelatinized tissue culture dishes in

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appropriate medium, at a sufficient dilution to result in individual colonies. Approximately 200 of these colonies are then picked for further analysis.

2) Transfection with plasmids containing the FLP 5 gene:

A plasmid containing the FLP gene under control of a promoter able to drive FLP expression, e.g., the human interferon-inducible 6-16 promoter, is constructed according to standard methods. Porter et al., EMBO J. 7: 10 85 (1988). Approximately 10 μg of FLP expression plasmid is transfected into approximately 10⁷ cells using standard electroporation conditions. With a plasmid containing the human 6-16 promoter, interferon is added at approximately 500 units/ml, in order to induce 15 expression of FLP. The cells are then treated as in (1), above.

The procedure to knock out the α -1,3-GalT gene in ES cells using an FRT-containing construct is:

- a) electroporate the complete construct into ES20 cells
 - b) select neo^R cells, and identify those ES cells having an interrupted α -1,3-GalT gene
- c) delete the neo^R gene using FLP recombinase, as described above; cells are tested for the excision 25 event as follows:

First, samples of each selected cell line are tested for the absence of the neo^R gene by treatment with the chemical G418. The cells will die in the presence of approximately 200 μg/ml G418 unless the neo^R gene is still present in the genome. Cell lines that are G418 sensitive are then tested further to confirm that excision of neo^R has occurred. This is done by Southern analysis or PCR analysis, both described in Sambrook et al. (1989). For Southern analysis, genomic DNA is isolated from the cells, digested with an appropriate

restriction enzyme, subjected to agarose gel
electrophoresis, and the digested DNA transferred to a
membrane. The DNA is hybridized with a labeled probe,
the label is detected (e.g., with X-ray film or color
development), and the pattern of bands indicates whether
or not an excision event had occurred in the cell line.
For PCR analysis, genomic DNA is isolated from the cells
and subjected to PCR reaction with suitable
oligonucleotide primers.

d) following confirmation of neo^R excision, the manipulated ES cells or PGC's are used to generate chimeric animals.

EXAMPLE 9

Preparation of DNA Constructs to Interrupt the α -1,3-GalT 15 Gene in Mice

Gene targeting (homologous recombination) is more efficient if the cloned cDNA fragments used for targeting are isolated from the cell line which is used for the gene knockout (i.e., the DNA is "isogeneic").

- 20 Accordingly, DNA was isolated from the E14 ES cell line (Hooper et al., Nature 326: 292-95 (1987)) and used to construct a mouse genomic library. The DNA was digested partially with the restriction enzyme Sau 3A, and fragments 12 kb 20 kb in size were isolated by glycerol 25 gradient fractionation. The size-fractionated DNA was
 - ligated into the Bam H1 site of \(\lambda\) EMBL3 (Sambrook et al. 1989, supra), and packaged in vitro to form lambda phage particles. The lambda library was plated by infection of E. coli strain PMC103 host cells (Doherty et al., Gene
- 30 124: 29-35 (1993)) at a density of 4x10⁴ phage per plate. A bovine cDNA clone, about 900 bp in length and containing a portion of the α-1,3-GalT gene corresponding to exons 7 9, was used to probe a total of 5.6x10⁵ independent recombinant phage. Four overlapping clones

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containing α-1,3-GalT gene sequences were isolated and
purified. The Sall restriction sites within these clones
were mapped (Figure 6), and the 4.0kb, 5.5kb, 11kb and
12kb SalI fragments from two of the clones (λ3 and λ5)

were subcloned into pBlueScript KS+ (Stratagene) or pUBS
(pUC19 carrying the pBlueScript KS+ polylinker) to
facilitate further detailed mapping of restriction sites.

These four subclones (designated pagt-S4.0, pagt-S5.5, pagt-S11 and pagt-S13) were mapped for restriction sites with restriction enzymes BamHI, EcoRI, HindIII, XbaI, XhoI, KpnI, SacI, SacII, EcoRV, PstI, SmaI, NotI and BglII. pagt-S4.0 and pagt-S5.5 were also checked for PvuI, PvuII, NdeI and SphI restriction sites. Detailed restriction maps of the 4 subclones were drawn from these data (Figures 7-12).

On the basis of these maps a knockout strategy was conceived. Basically the strategy is to insert a resistance gene (either neo^R or hyg^R) into the SalI site which lies within Exon 9. The knockout construct carries 20 the 4.0 and 5.5kb Sall fragments from p α GT-S4.0 and $p\alpha GT-S5.5$ which flank the Exon 9 SalI site (Figure 13). Screening for homologous recombination events then can be carried out using a DNA fragment representing the genomic region but lying outside the DNA included in the knockout 25 construct, i.e., outside the 9.5kb covered by $p\alpha GT-S4.0$ and pagt-s5.5. A 0.7kb EcoR1/XmnI fragment from pagt-s11 is used to screen Southern blots of BglII digested ES cell DNA for homologous recombinant events. band should appear on these Southerns when the 30 uninterrupted α 1,3-GalT gene is probed with this EcoR1/XmnI fragment (Figure 14). Insertion of the neo^R gene after a homologous recombination event will give rise to a 6.4kb band, due to the presence of a BglII site just flanking the Exon 9 SalI site within the knockout

construct. Thus the presence of the 6.4kb band is diagnostic for a homologous recombination event.

To carry out this strategy, the present inventors prepared a series of knockout constructs. The generation 5 of one such construct is outlined in detail in Figure 15. The vector paGT-S5.5, which carries the 5.5kb fragment immediately 3' to the Exon 9 SalI site, was chosen as the starting vector. paGT-S5.5 was digested with EcoRV and ClaI, generating a vector with a blunt end and a ClaI 10 compatible end. A 1:3kb fragment carrying the PMC1 promoter-driven neo^R gene flanked by FRT sites was excised from plasmid pNeo2FRT (previously constructed by the present inventors) by digesting with BamHI, filling in the restriction site and then digesting with ClaI to 15 generate a fragment with one blunt end and one ClaI compatible end. The nucleotide sequence of this 1.3kb fragment is provided in Figure 16 (SEQ ID NO: 13). fragment was then ligated into the ClaI/EcoRV digested pαGT-S5.5, the ligation mix transformed and colonies 20 screened for recombinants. One colony was recovered that contained the ${ t Neo}^{ t R}$ fragment inserted into the ${ t EcoRV/ClaI}$ of paGT-S5.5, based on the restriction pattern after digestion with diagnostic restriction enzymes ClaI, EcoRV, XbaI and EcoRI. This construct was designated 25 PNeoαGT6.8.

pNeoαGT6.8 was digested with SmaI, generating a vector with blunt ends. The 4.0kb Sall fragment was excised from pαGT-S4.0 and the ends filled. This fragment was then ligated into the SmaI digested pαGT-30 S5.5, the ligation mix transformed and colonies screened for recombinants. Four colonies were recovered which contained the 4.0kb SalI fragment inserted into the SmaI sites of pNeoαGT6.8 with the 5' portion of Exon 9 lying near the 3' portion of the exon in the nearby SalI 5.5kb fragment. The identity and orientation of the insert was

confirmed by the restriction pattern after digestion with diagnostic restriction enzymes XbaI, EcoRI, HindIII, BamHI, EcoRV and others. This construct was designated pNeo α GT10.8.

 $pNeo\alpha GT10.8$ was digested with ClaI, generating a 5 vector with ClaI compatible ends. Two complementary oligomers were synthesized that, when annealed, generated a linker containing translation termination codons in all three reading frames and a BglII site. The linker has 10 ClaI compatible ends. The linker was ligated into the ClaI digested pNeolphaGT10.8, the ligation mix transformed and colonies screened for recombinants. Many colonies were recovered that contained the linker inserted into the ClaI sites within $pNeo\alpha GT10.8$ based on the 15 restriction pattern after digestion with diagnostic restriction enzymes Bg1II, Cla and Bg1II/NotI. construct has been sequenced to confirm the identity, copy number and orientation of the insert. This construct is called pNeoaGT10.8B (Figure 17).

20

EXAMPLE 10

ES Cells - General Materials and Methods Working Conditions

Procedures for the isolation and culturing of all cell lines (embryonic stem, primordial germ and fetal fibroblast cell lines) require aseptic conditions to prevent growth of contaminating organisms:

- 1. All laboratory bench tops and equipment are wiped down with 70% ethanol prior to use.
- 2. All surgical instruments are autoclaved prior to use.
 - 3. Water for media preparation and cleaning of glassware is of high quality (e.g., Milli-Q water, prepared by passage through a Milli-Q ultrapure water system (Millipore).

- 4. Glassware is either dry-heat sterilized or autoclaved following extensive cleaning in Milli-Q water before use.
- 5. All tissue culture work is carried out under5 laminar flow conditions (Hepa filtered horizontal laminar flow workstation).
 - 6. All media are filter sterilized (22 μ m disposable filter) prior to use.
- 7. Antibiotics are used to minimize the risk of 10 bacterial contamination (Penicillin, Streptomycin and Gentamicin for bacteria; Nystatin for fungi).

Media/Solution Preparation

DULBECCOS MODIFIED EAGLE MEDIUM (DMEM)

10.0g DMEM powder- Gibco
15 (the low-glucose or high-glucose formulation, with or without pyruvate, may be used; L-glutamine is included)
1.0 liter Milli-Q-Water
3.7g NaHCO₃
Stir slowly until dissolved

20 Adjust pH ~ 7.2
Filter sterilize (following filter sterilization pH to rises to 7.4)
Keep at 4°C.

STO CELL MEDIUM

- 25 83.0 ml DMEM
 15.0 ml 15% fetal bovine serum (FBS); batch tested before use
 1.0 ml Pen/Strep 1:100
 1.0 ml Glutamine 1:100 (if needed) (see note below)
 - 30 Filter sterilize and keep at 4°C.

Note: Replenish complete medium (DMEM medium) (STO or ES) with glutamine.

*This step is only required if medium is older than 1 week - 10 days, as the glutamine breaks down after this 35 time.

ES CELL MEDIUM WITH OR WITHOUT LIF

up to 100.0 ml DMEM

- 68 -15.0 ml 15% FBS (batch tested before use; see below) 1.0 ml (from 0.01M stock) β -mercaptoethanol (0.1 mM final concentration) 5 1.0 ml Pen Strep. 1:100 0 - 1.0 ml Glutamine 1:100 (if needed) 1.0 ml Nystatin 1:100 0 - 2.5 ml Recombinant murine LIF (from 4x104 U/ml; 1000U/ml stock); activity-tested using LIF Assay 10 (see below) 0.4 ml Gentamicin 1.0 ml Nucleotides 1.0 ml Non-essential amino acids PENICILLIN/STREPTOMYCIN ANTIBIOTIC SOLUTION (1:100) 15 - Commonwealth Serum Laboratories, Australia; Catalogue No. 05081901 Penicillin G - 5000 U/ml

MITOMYCIN-C SOLUTION

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20 2.0 mg Mitomycin-C (Sigma Chemical Co. ("Sigma"); Catalogue No. M0503) 200.0 ml STO Cell Medium

Streptomycin Sulphate - 5000 μ g/ml.

Filter sterilize, divide into 20x 10 ml aliquot's and store at -20°C.

25 PHOSPHATE BUFFERED SALINE (PBS)

For 100 ml Milli-Q Water: (Ca⁺⁺ and Mg⁺⁺ - containing) (Ca⁺⁺ and Mg⁺⁺ - free)

	NaCl	0.89	0.80
	KC1	0.02	0.02
30	KH ₂ PO ₄	0.02	0.02
	$Na_2^2HPO_412H_2O$	0.289	1.115
	$CaCl_2 - 2H_2O$.014	-
	$MgCl_2 - 6H_2O$	0.01	• -
	Na pyruvate	0.0036	
35	D-glucose	0.1 g	-

Adjust to pH 7.4 and filter sterilize (Ca⁺⁺ and Mg⁺⁺ - free PBS is purchased from ICN Cell Biology and Tissue Culture, Cat. No. 18-604-54)

TRYPSIN/VERSENE (TV) WORKING SOLUTION (TV x 1)

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In PBS (Ca⁺⁺ and Mg⁺⁺ - free):
0.25% (w/v) trypsin (lyophilized)
0.04% (w/v) EDTA or EGTA

or:

5 To 1 liter of milli-Q water add the following:

		(Porcine, Difco)	2.5 g
	EDTA or EGTA		0.4 g
	NaCl		7.0 g
	$Na_2HPO_412H_2O$	mary angerous and the	$0.3 \bar{g}$
10	KH ₂ PO ₄	•	0.24 g
	KCl		0.37 g
	D-Glucose		1.0 g
	Tris		3.0 g
	Phenol red		1.0 ml

15 Adjust to pH 7.6, filter sterilize, aliquot and store frozen.

EGTA: Ethylene-glycol-bis(β -amino-ethyl ether)N,N,N',N'-tetra-acetic acid [Ethylene-bis(oxy-ethylenenitrilo)]tetraacetic acid

20 EDTA: Ethylenediaminetetraacetic Acid

Use either EDTA or EGTA. EGTA is preferred as it is less damaging to the ES/PGC cells.

GELATIN WORKING SOLUTION

0.1% gelatin in Milli-Q Water

25 Dissolve gelatin by heating to 60°C. Filter sterilize when still warm.

To gelatinize tissue culture plates:

- 1. Cover dish with solution, leave 30 minutes
- Aspirate gelatin and let dish air-dry.
- 30 NUCLEOSIDE STOCK SOLUTION

	Milli-Q Water	100 ml
	Adenosine (Sigma)	80 mg
	Guanosine (Sigma)	85 mg
	Cytidine (Sigma)	73 mg
35	Uridine (Sigma)	73 mg
	Thymidine (Sigma)	24 mg

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- 1. Dissolve by warming to 37°C.
- Filter sterilize and aliquot while warm.
- Store at 4°C or -20°C.
- 4. Thawing of nucleotides for use in ES cell media
 5 (a) nucleotides come out of solution upon thawing;
 - (b) Warm to 37°C to resolubilize before use.

NON-ESSENTIAL AMINO ACIDS (1:100)

- Commonwealth Serum Laboratories; Catalogue No. 09751301
- 10 100x concentrate for minimum essential medium (Eagle): (1.0 ml is added to 100 ml ES Cell Medium)

mg/10 ml milli-0 H20

	Glycine	7.5
	L-Alanine	8.9
15	L-Asparagine · H ₂ O	15.0
	L-Aspartic Acid [*]	13.3
	L-Glutamic Acid	14.7
	L-Proline	11.5
	L-Serine	10.5

20 WHITTEN'S CULTURE MEDIUM

	KCl	0.0356
	KH ₂ PO4	0.0162
	MgSO ₄ · 7H ₂ O	0.0294
	NaCl	0.4
25	NaHCO ₃	0.2106
	Glucose	0.1
	Na Pyruvate	0.0036
	Ca Lactate 5H ₂ O	0.0527
	Na Lactate	0.2416 ml
30	Milli-Q-H ₂ O	100 ml

The solution is adjusted to a final milliosmolarity of 250-280 by addition of $\rm H_2O$ or NaCl.

Filter sterilize and store at 4°C for two weeks.

Working solution:

35 10 ml

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Whitten's medium

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1.5g Diagnostic division, Code No. 81-001-4) BSA fraction V (Miles Pentex, Kankakee, Il., USA;

Filter Sterilize and equilibrate in $5\%0_2:5\%C0_2:90\%$ N₂ at 5.5%C, 95% humidity.

FBS BATCH TRIALS

Batches of FBS vary in the ability to support growth of ES cells, and in the ability to maintain the undifferentiated state of such cells. The following 10 procedure is used to identify suitable batches of FBS.

Use ES cells from between 2 & 20 passages:

Day 1

15

20

Split ES colonies and plate into dishes without feeder cells but with LIF. Incubate for 3 days.

Day 4 Trypsinise to detach colonies and cells.
Count cells and dispense into
gelatinized 6cm dishes
containing ES Cell Medium and
LIF (no serum added) as follows:

	Dish	Number	No. Cell	s	Batch	FBS Contro (Batch T	
	Non-Ina Serum	activated	·	A		B	esceu,
25	1	2	250	5	m1	-	_
	3	4	250	_		5 ml	_
	5	6	250	_		-	5 ml
	7	8	2000	5	ml	-	_
	9	10	2000	-		5 ml	_
30	11	12	2000	_		-	5 ml
	Inactiv	vated					
	Serum,	as control	(56°C for 15	min	.)		
	13	14	250	5	ml	-	_
	15	16	250	_		5 ml	_
35	17	18	250	_		_	5 ml
	19	20	2000	5	ml	-	-
	21	22	2000	_		5 ml	_
	23	24	2000	-		-	5 ml

There are duplicate plates for each treatment.

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Incubate low density dishes for 5 days Incubate high density dishes for 3 days

Day 7 Fix high density cells and stain with hematoxylin.

Day 9 Fix low density cells and stain for alkaline 5 phosphatase.

LIF ASSAY

This procedure is used to assay the potency of Leukaemic Inhibitory Factor (LIF).

Day 1 Split one 10 cm dish of confluent STO cells into five dishes. Incubate for 2 - 3 days in STO medium.

Day 3/4 When cells are confluent, replace medium with DMEM + 10% FBS. Incubate for 3 days.

Day 6/7 Collect conditioned medium (CM) and store at 4°C.

*Prepare low density ES cell cultures as described above.

	Dish	No.	Cells	C.M.	Med	ium	1000 LIF	U/ml	Medium w/o LIF	Presumed LIF Content
20	1,2,3	250		0.1 r	nl	4.9	ml	_	_	200 U/ml
	4,5,6	250		0.25	ml	4.75	ml	_	_	500 U/ml
	7,8,9	250		0.5	nl	4.5	ml	_	_	1000 U/ml
	10,11,12	250		1.0 r	nl	4.0		_		2000 U/ml
	13,14,15			_		_		5 ml	_	2000 0/1112
25	16,17,18	250		_		-		_	5 ml	

There are triplicate plates for each treatment.

Fix and stain for alkaline phosphatase.

Preparation of Fibroblast Feeder Cell Layers

Embryonic pluripotential cells are cultured in 30 vitro on a layer of fetal fibroblast cells. The fibroblast cells provide a wide range of factors necessary for the growth of pluripotential embryonic

cells (e.g. growth factors, cytokines, factors that are essential for maintenance of ES cell pluripotency).

ISOLATION OF PORCINE FETAL FIBROBLASTS:

- Remove developing porcine fetuses (preferably
 between days 16-30 of development) from uterus by aseptic dissection.
 - 2. Remove skin layer from fetus.
- Dissect out soft tissue avoiding developing wiscera.
 The white (fibroblast containing) tissue is found
 just under the skin layer.
 - 4. Wash dissected tissue in PBS (Ca⁺⁺ and Mg⁺⁺ free). Centrifuge at 1000 rpm for 5 min.
 - 5. Remove supernatant.
- 6. Incubate tissue in Trypsin/Versene Working Solution for 20 min.
 - 7. Dissociate cells by vigorously pipetting. Centrifuge at 1000 rpm for 5 min.
 - 8. Remove supernatant.
- Resuspend cells in STO Cell Medium. Allow large
 cell-clumps to settle.
 - 10. Plate out cells within supernatant (i.e., large cell clumps are not included) onto gelatinized tissue culture plates. Incubate cells in an atmosphere of 5% CO₂, 95% air (37.5°C, 95% humidity) until a
- confluent layer of fibroblast cells appears (~4-5 days).
 - 11. Passage of cells may be continued to increase cell numbers, or cells may be frozen or inactivated for further use.
- 30 CULTURE OF FETAL FIBROBLAST FEEDER LAYERS FROM FROZEN STOCKS:

Several different types of mouse feeder (STO cells) and porcine and bovine fetal fibroblasts can be used to form feeder layers. These include:

5

- (1) Bradley/Baylor mouse STO feeder cells that have been modified to express human LIF (gift from Allan Bradley, Institute for Molecular Genetics, Baylor College of Medicine, Texas Medical Center, Houston, Texas, USA)
- (2) Robertson/Columbia mouse STO feeder cells that have been modified to express murine LIF (gift from Elizabeth Robertson, Columbia University, New York, USA)
- 10 (3) Several porcine fetal fibroblast lines
 - (4) Several bovine fetal fibroblast lines

(the fibroblast lines of (3) and (4) were derived by the present inventors using the procedures described above)

- 15 The procedure for producing feeder layers is as follows:
 - 1. Rinse one 10 cm tissue culture (tissue cure) dish with gelatin/Milli-Q water solution for 30 min. Aspirate gelatin solution and let dish air-dry.
- 3. Add 10 ml of STO cell medium to 15 ml centrifuge 20 tube.
 - 4. Remove feeder layer cells frozen in freezing media from liquid N_2 container.
 - 5. Thaw cells by warming vial in hands or in 37°C water bath.
- 25 6. Transfer STO cells to medium in centrifuge tube.
 - 7. Spin at 1000 rpm for 5 min.
 - 8. Resuspend cells in 10 ml medium and transfer to gelatin-treated tissue culture dish.
 - 9. Incubate at 37°C for 3 days.
- 30 SPLITTING OF FEEDER LAYER STO CELL/FETAL FIBROBLASTS:

 This procedure is used to expand the number of cells from a single confluent plate/dish; cells are detached from the confluent plate and transferred to fresh plates at sub-confluent densities.
- 35 1. Gelatinize five 10 cm tissue culture dishes.
 - 2. Examine incubated STO cells under microscope and check for confluence.

- 3. If STO feeder monolayer is confluent (cells cover bottom of dish, or nearly so), wash gently with PBS (Ca⁺⁺ and Mg⁺⁺ free) for 1 min.
- 4. Aspirate PBS and add 1 ml Trypsin/Versene Working
 5 Solution for 1 min (or until cells start to detach).
 Check under microscope.
 - 5. Detach cells by vigorously pipetting, add 1.0 ml STO Cell medium (i.e., a ratio of 1:1 STO Cell medium:Trypsin/Versene Working Solution) to
- neutralize trypsin, and transfer to a centrifuge tube containing 10-15 ml STO Cell medium. Wash cells remaining on dish with some of STO cell medium from the tube. Centrifuge at 1000 rpm for 5 min., aspirate supernatant, resuspend pellet in 1 ml STO
- 15 Cell medium. Resuspend cells to make single cell suspension. Make up to 50 ml with STO Cell medium.
 - 6. Dispense 10 ml into each of the five tissue culture dishes and incubate until confluent (~ 3 days).

INACTIVATION OF FEEDER LAYERS:

The present inventors use two alternative methods for inactivating feeder layers, which stops the cells from dividing:

(1) Mitomycin treatment:

- Check dishes for confluence of STO cells/fetal
 fibroblasts.
 - 2. Thaw mitomycin-C solution and use undiluted.
 - 3. Aspirate STO cell medium from feeder cell plate.
 - 4. Add 10 ml aliquot of mitomycin-C to plate and incubate at 37°C for 1-3 hours.
- 30 5. Aspirate mitomycin-C, wash cells in 1x PBS (without Ca^{++} or Mg^{++}) for 1 min.
 - 6. Aspirate PBS and add 1 ml trypsin solution for 1 min.

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- 7. Detach cells by vigorously pipetting and transfer to STO cell medium in centrifuge tube.
- 8. Centrifuge at 1000rpm for 5 min.
- 9. Resuspend cell pellet in 1 ml ES Cell Medium.
- 5 10. Plate out in dishes in preparation for addition of ES cells.
 - (2) Gamma Irradiation:
 - 1. Check dishes for confluence of STO cells/fetal fibroblast.
- Trypsinise cells into single cell suspension.
 - Irradiate cells (3000 rads) in STO cell medium.
 - Centrifuge at 1000 rpm for 5 min.
 - 5. Resuspend pellet in 1 ml ES Cell Medium.
- 6. Transfer cells to gelatinized tissue culture dishes with ES Cell Medium and place in incubator at 37°C until the cells adhere to the dish. NOTE: If cells are not confluent, count using hemocytometer and seed at 5x10⁴ cells in 1 ml medium per well of Nunc 4-well plate.

One 10 cm dish of inactivated cells can be split into:

Ten 4-well plates (Nunc tissue culture plates), or Eight 3.5 cm tissue culture dishes, or Three 6 cm tissue culture dishes, or

Two 20 cm tissue culture dishes.

<u>Demonstration of Totipotency:</u>

A. <u>Blastocyst Injection</u>

The ability of embryonic cell lines to form germline chimeric animals is a conclusive test for their

30 totipotency. This can be accomplished by blastocyst injection experiments, using techniques for various mammalian species substantially the same as those established for the mouse. See Example 14, below. See also, e.g., Bradley, Production and Analysis of Chimeric

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Mice, In: <u>Teratocarcinomas and Embryonic Stem Cells: A</u>

<u>Practical Approach</u> (E.J. Robertson, ed.), IRL Press,
Oxford, pp. 113-52 (1987). However, for porcine
manipulations the holding pipette must be somewhat larger
as porcine embryos are larger than mouse embryos.

B. <u>Co-Culture of ES Cells/PGC's and Morula Embryos</u>

Embryos at the morula stage of development are surgically collected from superovulated animals. For porcine embryos, for example, the zona pellucida is then disrupted using Acid Tyrodes solution and ES cells/PGC's are cultured in the presence of the zona pellucidadisrupted morulae. ES/PGC cells adhere to the exposed morula cells and, following overnight culture in Whitten's medium, the embryos are transferred to synchronized recipients. Preferably, the zona pellucidadisrupted morula is completely free of the zona pellucida. However, this need not be the case as long as the ES cells/PGC's can gain direct access to at least some of the morula cells.

20 C. Morula Injection

embryo prior to formation of the blastocyst cavity. The technique is similar to blastocyst injection. Es cells or PGC's are drawn into an injection pipette, which is inserted beneath the zona pellucida. Then, the cells are expelled so that they are in contact with the cells of the morula embryo. The injected morula is then cultured overnight in Whitten's medium (porcine) or other appropriate medium to allow blastocyst formation.

BNSDOCID: <WO____ 9520661A1 | >

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D. <u>Nuclear Transfer and Embryo Cloning</u>

ES cells and PGC's can be fused to enucleated zygotes that have been derived by in vitro maturation, in vitro culture, in vitro fertilization or collected 5 surgically. Following successful fusion the embryos can be transferred to synchronized recipients. In vitro or in vivo-collected porcine oocytes, for example, are manipulated in Whitten's medium supplemented with 1.5% BSA Fraction V and 7 μ g/ml cytochalasin B (Sigma). 10 bevelled micropipette is used to remove the metaphase plate from the oocyte. A single ES cell or PGC (after trypsin treatment to form a single-cell suspension) is inserted through the zona using a bevelled micropipette, such that the cell comes in contact with the oocyte 15 plasma membrane. Fusion is achieved in a 28 V/cm AC field for 5 sec. followed by an 80 V/cm DC pulse of 100 μ sec. duration. Subsequent to observed fusion, embryos are incubated at 39° C in 5% CO_2 , 5% O_2 , 90% N_2 in microdrops of Whitten's medium supplemented with 1.5% 20 BSA, until transfer to a synchronized recipient.

EXAMPLE 11

Murine ES Cell Culture

ES cells are able to differentiate spontaneously into many different cell types, and culture conditions

25 which prevent this differentiation are critical for the continuous passage of these cells in an undifferentiated form, capable of contribution to chimeric mice.

I. CULTURE CONDITIONS

ES cells are grown in polystyrene cell culture
30 dishes treated with 0.1% gelatin (made up in PBS or
Milli-Q water) for 10 minutes. A feeder layer of
mitotically inactivated fibroblasts provides a source of
cytokines. The fibroblasts are either primary mouse
embryo fibroblasts (PMEFs), or STO fibroblasts, an

immortal line. The medium used is DMEM supplemented with glucose, amino acids and nucleosides. Robertson, Embryo-Derived Stem Cell Lines. <u>In</u>: Teratocarcinomas and Embryonic Stem Cells: A Practical Approach (E.J.

5 Robertson, ed.), IRL Press, Oxford (1987). To this medium is added LIF (final concentration of 10³U/ml Esgro, AMRAD). FBS is added to 15%. The batch of FBS is chosen on the basis of its ability to support ES cell growth with low levels of differentiation (i.e, only rare individual cells undergo differentiation. The ES cells are grown in an atmosphere of 5-10% CO₂, at 37°C II. ROUTINE PASSAGE

ES cells must be passaged frequently to prevent the colonies from growing too large and differentiating.

15 This is achieved by splitting the cells at a ratio of 1:10 to 1:40, every two to four days.

EXAMPLE 12

Genetic Manipulation of Cells

The general procedures set out in this Example

20 provide guidelines that are readily adaptable to
 individual experimental situations that might employ, for
 example, different cell lines or equipment supplied by
 different manufacturers. This Example also provides
 specific procedures used and results obtained in

25 generating a set of mouse ES cell lines in which the α 1 3 galactosyltransferase gene was disrupted by homologous
 recombination. The general procedures provided in this
 Example are adapted for mouse ES cells. However, the
 procedures are substantially similar for porcine ES

30 cells.

- I. INTRODUCTION OF DNA INTO ES CELLS BY ELECTROPORATION
- A. Coat required number of plates with 0.1% gelatin (in PBS or Milli-Q water). (Usually 2 X 6 well plates and 8 well plate)
- B. Thaw 10⁷ embryonic fibroblasts into DMEES (equivalent to ES Cell Medium); inactivate by irradiating at 3000 Rad.
 - C. Count irradiated cells, spin down and resuspend in DMEES to 10^6 cells/ml.
- D. Aspirate gelatin from plates and plate cells at: 7 X 10⁵ cells/well (6 well plate) in 2.5ml medium; 7 X 10⁴ cells/well (24 well plate) in 1 ml medium.

 Incubate at 37°C, 5-10% CO₂ for 3 4 hr.
- E. Wash ES cells in 5 ml (250 ml flask) PBS-EGTA and let sit at room temperature for 4 min.
 - F. Remove PBS, add 5 ml trypsin (CSL) and leave at room temperature for 2-4 min. Wash down cells, add 10 ml DMEES and count. Approximately 5 X 10^6 to 2 X 10^7 ES cells are needed for experiments.
- G. Centrifuge cells and resuspend in 10 ml PBS. Centrifuge again and resuspend in 540 μ l PBS. Dilute 50 μ l into 10 ml DMEES and culture to determine plating efficiency.
- H. Add 5 10 μ g DNA to cells in 10 ul PBS (total 25 volume, 500 μ l) and transfer to sterile electroporation cuvette (e.g. Biorad).
- I. Electroporate at 0.22 kV, 500 μ FD (time constant should be ~8.4). This is achieved using a Biorad Gene Pulser unit (Biorad Catalogue No. 1652078) with capacitance extender (Biorad Catalogue No. 1652087), or similar device.
 - J. Resuspend in 10 ml DMEES with constant pipetting to break up clumps of DNA from lysed cells.
 - K. Centrifuge cells and resuspend in 5ml DMEES.

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L. Take 50 μ l, add 50 μ l trypan blue solution and count for viability.

M. Culture by dilution plating to determine plating efficiency.

5 II. SELECTION CONDITIONS

ES cells that do not express a neomycin resistance gene are selectively killed by treatment with G418 at 200-500 µg per ml of medium. Antibiotic- containing medium is changed daily. A population of cells that has not been electroporated also is treated in order to see how genuinely sensitive cells respond to the G418 treatment. After 6 to 10 days, cells resistant to the antibiotic will be evident as healthy colonies. These cells will have been transformed by the targeting construct and can be screened for homologous recombination (i.e., screened for gene targeting versus random integration).

Resistant colonies are picked from the selection dish with a mouth pipette and dispersed into a single cell suspension. Half of these cells are frozen away while the other half is expanded and used to determine whether or not homologous recombination has occurred. In the colonies are small, it is sometimes preferable to expand the whole colony in a 24 well dish, and then to freeze half while further expanding the other half for genetic analysis.

III. PICKING ES CELL COLONIES FOR GENETIC ANALYSIS
AFTER SELECTION

A. <u>Method 1: Freezing Half Colonies</u>

The day before colony picking:

a) Coat required number of plates with 0.1% gelatin (in PBS). Two plates per 24 colonies to be picked: one plate is for freezing and one plate

is for clone expansion. Start with 20 X 24 well plates.

- b) Count irradiated fibroblasts, spin down and resuspend in DMEES.
- c) Aspirate gelatin from 10 plates and plate ~10⁵ (can use as few as 5 X 10⁴) cells/well in 1ml DMEES. Incubate at 37°C, 10% CO₂ overnight (or a minimum of 1 h).
- d) Aspirate the gelatin from the other 10 plates.
- On the day of colony picking:

 a) Change medium on ES cells before and regularly during picking (to remove floating cells).
 - b) Pull plugged pasteur pipettes. Use a fresh pipette after each 24 colonies. The desired tip is about half a colony in diameter, with the constriction over 1-2cm. The tip should be perpendicular and neat. Note: after drawing the pipette, rub the glass at the desired break point with freshly drawn glass, then bend.)
 - c) Label multi-tip reservoirs for:
 1 PBS-EGTA
 2 Trypsin-Versene
 3 DMEES
 4 2 X Freezing mix(20% DMSO in FCS)
 - d) Using multipipettor, dispense 50 μ l PBS-EGTA into 24 wells of 96 well plate.
 - e) At microscope: Connect finely drawn pasteur pipette to mouth pipette tube. Dislodge colony from plate and transfer (in minimum volume) to one well of a 96 well plate. Expel contents of pipette; the bubbles serve as a location guide. Pick 24 colonies or as many as possible in <10-15 min (preferably a multiple of 6).

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		f)	Back in hood: Add 100 μ l trypsin to each well using multipipettor) and leave at RT for 2 min.
5		g)	Pipette up and down 10- 15X to disperse cells, then add 100 μ l DMEES. (This should be done within 4-6 min after trypsin addition).
10	of the state of th	· i)	Divide cell suspension between freezing and expansion plates using 12 channel pipette with every second tip fitted. Transfer 125 μ l to gelatinized 24 well plate (to freeze); the remaining ~125 μ l is
15			transferred to a 24 well plate with feeder layer (for DNA). The plates are labelled and carefully aligned to ensure that one clone goes into the same well of each tray.
20		j)	Add 125 μ l 2 X freeze mix to each well on freezing plate, mix well by swirling.
25		k)	Seal in ziplock bag or plastic wrap and place in -70°C freezer in an equilibrated styrofoam box. Interleave the plates with styrofoam sheet.
		1)	Incubate expansion plates until there are sufficient cells for genotype analysis.
30	A. <u>M</u>	ethod 2:	Freezing after expansion to 24
	<u>w</u>	ells.	
	1	. The	day before colony picking:
35		a)	Coat required number of plates with 0.1% gelatin (in PBS). Start with 10 X 24 well plates.
		b)	Count irradiated fibroblasts, spin down and resuspend in DMEES.
40		c)	Aspirate gelatin from the plates and plate ~10 ⁵ cells/well in 1ml DMEES. Incubate at 37°C, 10% CO ₂ overnight (or a minimum of 1 h).

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On the day of colony picking	2.	On	the	day	of	colony	picking
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- a) Pick colonies as described for half colonies (method 1, above) but instead of dividing the cell suspension between freezing and expansion plates, the entire cell suspension goes into the expansion plate.
- b) After 3-4 days (with daily medium changes) the cells will have grown sufficiently to be frozen. Working one plate at a time (with practice two can be handled), aspirate medium from each well. Flood with PBS/EGTA for 4 minutes. Meanwhile, set up pipette tips to fit alternate channels of a twelve channel multipipettor. Aspirate PBS.
- c) Add 100 μ l trypsin (using multipipettor and alternate channels) and leave at room temp. for 2 min.
- d) Pipette up and down 10- 15X to disperse cells of first row, change tips, then add 100 μ l DMEES. Repeat for each row. (This should be done within 6 min of trypsin addition).
- e) Using 12 channel pipette with every second tip fitted, transfer 125 μ l to gelatinized 24 well plate (to freeze). The remaining cells will be expanded for DNA. It is crucial that the plates are labelled and carefully aligned to ensure that the freezing tray matches the expansion tray.
- f) Add 125 μ l 2 X freeze mix to each well on freezing plate; mix well by swirling.
- g) Seal in ziplock bag or plastic wrap and place in -70°C freezer in an equilibrated styrofoam box. Interleave plates with styrofoam sheets.
- h) Add 1ml of DMEES to the expansion tray. (There will be sufficient feeder cells to give good plating

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efficiency). Incubate for 3-4 days until there are sufficient cells for genotype analysis.

IV. THAWING OF ES CELL CLONES FROZEN IN 24-WELL PLATES

Cells that have been identified to have the desired genetic alteration are recovered from a duplicate plate frozen at -70°C. The plate is taken to the laminar flow hood and removed from the plastic bag. Each well is filled with warm medium, and feeder cells are added to the well(s) of interest. The plate is placed in a 37°C incubator for 60 min., then the medium is replaced. Colonies will appear after two or three days. These colonies are expanded for establishment of new frozen stocks, and tested for 1) karyotype analysis; 2)

confirmation of the desired genetic alteration; 3) mycoplasma infection; and 4) ability to form chimeras.

EXAMPLE 13

Production Of Mouse ES Cell Knockouts Using The pNEOαGT10.8B Construct

20 I. TRANSFORMATION

A total of 1×10^7 E14 ES cells was electroporated with 5μ l of $1\mu g/\mu$ l pNeo α GT10.8B DNA (linearized by XhoI digestion) (see Example 9 and Figure 17). Electroporation was carried out in 600μ l in a wide cuvette at 25μ F, 350V for 0.5msec. Cells were recovered in 6ml ES complete

medium and plated into 6 x 100mm petri dishes, each containing a feeder layer of Neo^R STO cells.

Cells were cultured in ES complete medium for 3 days and then medium containing 200-350 μ g/ml G418 was 30 substituted. This medium was changed every second day. After 9 days, individual Neo^R colonies were sufficiently large to be identified and recovered. Colonies were picked in 20 μ l PBS and 20 μ l of trypsin solution were added. Forty μ l of 60% BRL conditioned medium in ES

complete medium were then added. Aliquots of $40\mu l$ were transferred to single wells of each of two 24-well plates. One plate contained a feeder layer of STO cells in $100\mu l$ ES complete medium. $140\mu l$ of 2x DMSO freezing mix was added to this plate, which was stored at -80°C . Each of the wells of the second 24-well plate contained 1ml of 60% BRL conditioned medium in ES complete medium. This plate was incubated at 37°C until the colonies were confluent.

- 10 II. CONFIRMATION OF HOMOLOGOUS RECOMBINATION
 Medium was aspirated off confluent colonies and
 400μl lysis buffer (10mM Tris pH 7.8, 100mM NaCl, 1mM
 EDTA, 1% SDS, and 500μg/ml Proteinase K) added. The
 cells were lysed at 37°C overnight, extracted with 400μl
 15 1:1 phenol/chloroform and transferred to Eppendorf tubes
 containing 1ml 95% ethanol and 0.2M NaAc. DNA was
 pelleted by centrifuging at 13,000 rpm in an Eppendorf
 centrifuge, the pellet washed twice with 80% ethanol and
 redissolved in 30μl water.
- Southern analysis (see, e.g., Sambrook et al., supra) was used to identify ES cell clones where homologous recombination had occurred at the 3' end of the construct. Aliquots of 15µl of DNA were digested with 20 units of the restriction enzyme BglII according to the manufacturer's recommendations. After incubation at 37°C overnight, the DNA was electrophoresed through a 0.8% agarose gel (in a Tris acetate, EDTA buffer) at 1-2V/cm overnight, using 750ng of HindIII-digested lambda DNA as markers. The DNA was transferred to a Zetaprobe nylon membrane using a Hybaid vacublotter at a vacuum of 80cm Hg for 1 hour.

The membrane was prehybridised in a Hybaid hybridization bottle in 10ml of the following hybridization mix for 3 hours at 65°C:

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0.25M Na₂HPO₄ pH 7.2 7% SDS 1mM EDTA 100 μ g/ml salmon sperm DNA 10% PEG

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Radioactively labeled probe DNA was prepared using a BRESATEC gigaprime oligo labeling kit (Cat. No. GPK-1) according to the manufacturer's recommendations. Approximately 50ng of a 0.7kb EcoRI/XmnI DNA fragment from beyond the 3' terminus of the construct pNeoαGT10.8B (see Example 9 and Figure 17) were labeled with ³²P-dATP to a specific activity of 5x10⁸ cpm/μg. The denatured probe was added to the prehybridising membrane in the Hybaid bottle and incubated overnight at 65°C.

The membrane was removed from the Hybaid bottle, rinsed with 0.5xSSC, 0.1% SDS prewarmed to 65°C, and then washed 2-3 times with 0.1xSSC, 0.1% SDS at 65°C for 30 min each wash. Excess moisture was then blotted from the membrane, the membrane wrapped in plastic wrap and exposed to a phospho-imager screen for 16 hours up to 3 days. The image was visualized on an Imagequant phospho-imager.

Results are shown in Figure 18, which is a Southern blot of DNA from 15 ES cell lines probed with the 25 diagnostic 0.7kb EcoRI/XmnI DNA fragment described above and in Example 9. The 6.4kb band, diagnostic for a homologous recombination event in the α 1-3 galactosyltransferase gene (α 1-3 Gal T) (see Example 9), is seen in 6 of the 15 ES cell lines examined. All of the 6 knockout cell lines appeared to be heterozygous for the inactivated allele since the 8.3kb band, diagnostic for the uninterrupted α -1,3-Gal T gene (see Example 9), was also present in all six lanes.

Two cell lines, designated hereinafter "8D1" and 35 "7C2," were chosen for further analysis. Cell lines 8D1 and 7C2 were identified by Southern analysis to contain

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an α -1,3-Gal T allele where homologous recombination had occurred at the 3' boundary of the construct.

Long range PCR was then used to determine whether or not homologous recombination had occurred at the 5' 5 boundary of the construct within these cell lines. Two sets of primers were used in separate PCR experiments:

1) Wild-type primers:-

MGT-KOex8F and MGT-KOR1 span the intron between exons 8 & 9, and amplify a 5.5 kb fragment from the wild10 type α-1,3-GalT gene (Figure 19)
SEQUENCES:

MGT-KOex8F

5'TGCTGGAAAAGTACTACGCCACACAGAAACTCA-3'

(SEQ ID NO: 14)

15 (Nucleotides 1014-1046 in Figure 4)

MGT-KOR1

5'AGCCAGAGTAATAGTGTCAAGTTTCCATCACAA-3'

(SEQ ID NO: 15)

(Nucleotides 1779-1811 in Figure 4)

20 2) Knockout primers:-

MGT-KOex8F and MGT-KONeoR span exon 8 to the Neo^R gene cassette in the "knock-out" allele and amplify a 5.5 kb fragment from the knocked out allele (Figure 19) SEQUENCE:

25 MGT-KONeoR

5'-GCCACACGCGTCACCTTAATATGCCAAGTGGAC-3'

(SEQ ID NO: 16)

(Nucleotides 323-355; Figure 16)

Each reaction contained ~100 ng genomic DNA as template in a reaction volume of 50μl and contained 25mM Tris HCl (pH9.1), 16mM (NH₄)₂SO₄, 250 μM dNTPs, 3.5 mM MgCl₂, 100 ng each primer, 2 units Taq polymerase and 0.025 units Pfu polymerase. The reactions were heated at 94°C for 1 min, then 45 cycles of 94°C for 15 sec, 68°C

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for 6 min, followed by a single step of 72°C for 10 min. Genomic DNAs from putative "knock-out" ES cell lines from CBA/C mice (homozygous for the wild-type α -1,3-Gal T allele) were amplified in separate reactions using each set of primers. A $10\mu l$ aliquot of each PCR was analyzed by Southern blotting (Sambrook et al., 1989).

The results are illustrated in Figure 20: Knockout primers:-

A 5.5 kb fragment that hybridized to the 1.3 kb Neo^R
gene cassette (Figure 16) was generated from 7C2 DNA
(Figure 20; lane 4) and 8D1 DNA (not shown). This
band was not generated from CBA/cDNA (Figure 20;
lane 3).

Wild-type primers:-

A 5.5 kb fragment that hybridized to the α-1,3-Gal T gene probe (isolated by Sal I digestion of pαGT-S4.0) was generated from 7C2 and CBA/cDNA's (Figure 20; lanes 1 and 2 respectively) and 8D1 DNA (not shown). This product did not hybridize to the Neo^R gene 20 probe.

These results demonstrate that homologous recombination had occurred at the 5' boundary of the construct in cell lines 8D1 & 7C2.

EXAMPLE 14

- 25 <u>Generation of Animals Carrying an ES Cell Genome</u>

 The procedures provided in this Example are adapted for mouse ES cells. However, the general strategy is substantially the same for porcine ES cells and PGC's.
- I. PREPARATION OF ES CELLS FOR INJECTION

 ES cells are split into wells of a 24-well dish at cell densities of 1:2, 1:4, 1:8 and 1:16, relative to the initial density, two and three days before injection.

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The most vigorous and least differentiated cultures are chosen on the basis of morphology.

II. EMBRYO INJECTION AND PRODUCTION OF CHIMERIC MICE

Mouse embryos are collected from either superovulated or naturally mated female mice, approximately 3.5 days after mating. After overnight culture in M16 medium (Bradley, Production and Analysis of Chimaeras. In Teratocarcinomas and Embryonic Stem Cells a Practical Approach (E.J. Robertson, ed.) IRL Press, Oxford, pp. 113-10 52 (1987)), those that have cavitated to form blastocysts are microinjected with about 12 to 20 ES cells. This microsurgical procedure is performed with instruments drawn from capillary glass, and injection is controlled with micrometer syringe-based hydraulic devices. A differential interference contrast-equipped inverted microscope is used to view the procedure.

After injection, blastocysts are transferred to the uterus of pseudopregnant female mice. Chimeric mice are identified by coat color contribution by the ES cells.

20 Chimaeric mice show agouti coat colour derived from the host blastocyst, and chinchilla contributed by the ES cells.

Chimeric mice were generated from ES cells carrying the interrupted a-1,3-Gal T allele (including 8D1, 7C2 cells) by injection into C57B1/6J x CBA F2 blastocysts. The ability of individual chimaeric mice to transmit the ES cell characteristics through the germ-line was estimated by glucose phosphate isomerase (Gpi) analysis of sperm (Bradley, supra, (1987)); Mann et al., J. Reprod & Fert. 30 99, 505-512 (1993). Glucose phosphate isomerase catalyses the interconversion of glucose-6-phosphate to fructose-6-phosphate. Mice have a single structural Gpi locus with two main alleles Gpi 1A and Gpi 1B. Gpi 1A codes for protein which appears as a slow cathodically migrating band

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during electrophoresis and occurs in strains such as BALB/c and C129. (The ES cells used here were derived from strain 129 mice). Gpi 1B determines an enzyme that moves faster than Gpi 1A and occurs in the wild and in strains such as 5 C57 and CBA (used here to derive host blastocysts).

Heterozygotes have the two parental bands plus an intermediate band which indicates the dimeric structure of the enzyme. Multiple electrophoretic forms occasionally observed are due to oxidation of sulfyhdryl groups and not due to tissue-specific expression. In chimaeric mice, the ratio of Gpi 1A (strain 129-derived) to Gpi 1B (derived from the host blastocyst) indicates the proportion of cells with the ES cell genotype within different tissues. The appearance of Gpi 1A (derived from the ES cells) in the 15 sperm suggests that the mouse is able to transmit the ES cell genotype through the germ-line.

III. GENERATION OF MICE HOMOZYGOUS FOR THE GENETIC CHANGE INTRODUCED INTO THE ES CELLS.

Chimaeric mice with sperm derived from ES cells were 20 mated to BALB/c mice. Offspring with the 129/Ola X BALB/c genotype (i.e. heterozygous for the ES cell genotype) are grey. Half of these grey mice were expected to carry the interrupted allele. Mice heterozygous for the interrupted allele were identified by PCR analysis of genomic DNA obtained from blood.

To generate mice homozygous for the inactivated $\alpha-1,3-$ Gal T gene, the heterozygous mice were mated to each other. One quarter of the offspring were expected to be homozygous for the interrupted gene. Homozygotes were identified by PCR analysis of genomic DNA obtained from blood. The PCR strategy was based on the insertion of a Neo^R gene in the Sal I site of exon 9 of the $\alpha-1,3-$ Gal T gene (Figure 13). Wild-type primers:-

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E9F: 5'TCAGCATGATGCGCATGAAGAC 3'

(SEQ ID NO: 17)

(homologous to sequence about 40 to 60 bp 5' to the Sal I site of exon 9, corresponding to nucleotides 1257-5 1278; Figure 4)

E9R2: 5'TGGCCGCGTGGTAGTAAAA 3'

(SEQ ID NO: 18)

(homologous to a region about 175 to 195 bp 3' to the Sal I site of exon 9, corresponding to nucleotides 1511-10 1492; Figure 4)

The expected fragment size generated from the wildtype allele is 255 bp (Figure 21). These primers also can
potentially generate a 1596 bp PCR fragment from the
interrupted allele. In practice this fragment was not
generated when both the wild-type and interrupted alleles
were present, probably because the smaller 255 bp product
is amplified preferentially.

Knock-out primers:-

NeoF1: 5' TCTTGACGAGTTCTTCTGAG 3'

20 (SEQ ID NO: 19)

(corresponding to nucleotides 1170-1189; Figure 16)

E9R2: (the same primer described above to detect the wild-type allele)

The expected fragment size is 364 bp (Figure 21).

Mice were grown to weaning age and bled from the tail. Sodium Heparin was added to about 10 U/ml. PCR amplification was conducted on 1 μ l of heparinised blood (~10⁴ nucleated cells) in a 50 μ l reaction volume containing 100 mM Tris-Acetate pH 8.8, 3.5 mM MgCl₂, 0.2mM dNTPs, and 2 units Tth DNA polymerase. Each reaction contained both the wild-type and knock-out primers at a concentration of 2ng/ μ l for each primer. To ensure that Tth polymerase was

not inhibited by heparinized blood, each reaction was performed in duplicate.

One of the reactions was spiked with two DNA samples:

- i) 10 fg (~600 molecules) of linearized KO plasmid 5 pNeo α GT10.8B.
 - ii) 1 fg (~1000 molecules) of a 983 bp RT-PCR product that includes Exon 9.

The other reaction was not spiked. Thus, two separate PCR reactions were set up for each blood sample. In addition, 10 control PCR reactions with no genomic DNA template and with or without spikes were conducted. Each reaction mix was heated at 94°C for 3 min., then incubated for 40 cycles at 94°C for 40 sec., 53°C for 40 sec., and 72°C for 40 sec. Aliquots of 5 µl of each reaction mix were electrophoresed on a 3% agarose gel, and DNA fragments were visualized on a UV light box after staining with ethidium bromide. HpaII-digested pUC19 plasmid DNA was used for markers.

Results of the PCR analysis for three mice, and a "no DNA" control, are shown in Figure 22. For mouse #42, the 20 KO primers generated a 364 bp band in the + spike reaction only. The wild-type primers generated a 255 bp band in the + spike and - spike reactions. These results demonstrate that mouse #42 is homozygous for the wild-type allele. For mouse #43, the wild-type primers generated a 255 bp band in 25 the + spike reaction only. The KO primers generated a 364 bp band in the + spike and - spike reactions. results demonstrate that mouse #43 is homozygous for the interrupted allele. For mouse #44, the KO generated a 364 bp band in the + spike and - spike The wild-type primers generated a 255 bp band in the + spike and - spike reactions. These results demonstrate that mouse #44 is heterozygous In the control PCR reactions, no interrupted allele. product was evident when template was not included. PCR 35 products of 364 bp and 255 bp were evident

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pNeoαGT10.8B and Exon 9 RT-PCR DNA were the only templates included in the control reactions.

EXAMPLE 15

Characterization of Homozygous Knockout Mice

5 I. ABSENCE OF Gal T mRNA IN Gal T KNOCKOUT MICE

A. RNA Isolation

Total RNA was extracted using the RNAzol™B kit (BIOTECX Laboratories, Inc., 6023 South Loop East, Houston, Texas 77033, USA.), supplied by Bresatec. This extraction 10 procedure is based on the method described by Chomczynski et al., Anal. Biochem. 162: 156-159 (1987), and involves homogenization in a guanidinium/phenol solution, chloroform extraction, 2 isopropanol precipitations, and 75% EtOH washes. The RNA was stored as an EtOH precipitate 15 at -20°C and quantitated by measuring absorption at wavelenth 260 nm in water. The integrity and quantitation was confirmed by electrophoresis in agarose/formaldehyde gels. Sambrook et al. Molecular Cloning. A Laboratory Manual. Second Edition. (1989)

B. RT-PCR

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First strand cDNA synthesis involved:

- annealing $2\mu g$ of total RNA from kidney, heart or liver with 120ng oligo dT primer (Gibco BRL, M-MLV Reverse Transcriptase Kit) at 65°C for 5 minutes in $5\mu l$ of 10 mM Tris-HCl,1mM EDTA (pH8).
- reverse transcription at 37°C for 1-2 hours in a final reaction volume of $20\mu l$ utilizing the M-MLV Reverse Transcriptase Kit(Gibco BRL). Each reaction contained 5mM DTT, $0.1\mu g/\mu l$ BSA, 1mM dNTPS, 40 U of human placental RNAse Inhibitor (Bresatec), 200U of M-MLV Reverse Transcriptase and the associated RTase buffer at 1X concentration.

C. PCR Analysis of cDNA

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 \mathbf{T} CDNA α -1,3-Gal was detected by PCR amplification of oligo dT-primed cDNA template. Failure to generate this PCR fragment, in conjunction with the control PCR results, indicated that $\alpha-1,3$ -Gal T mRNA was absent from the RNA preparation. demonstrate that the $\alpha-1,3$ -Gal T primers supported amplification of the $\alpha-1,3$ -Gal T template, reaction was assembled in duplicate, and one of the reactions was spiked with 0.1 fg (~100 molecules) of a 983 bp mouse $\alpha-1,3$ -Gal T cDNA product (generated by primers 7F and mGT-3UR, spanning exon 7 to the 3' untranslated region). As a second control demonstrate that cDNA synthesis had occurred, ferrochelatase PCR fragment was generated from the cDNA template.

1. Primers:

Primers to detect α -1,3-Gal T cDNA:

7F: 5'- TGGAGATCGCATTGAAGAGC 3'

(SEQ ID NO: 20)

20 (corresponding to nucleotides 889-911

within exon 7 (Figure 4)

9R2: 5'- TGGCCGCGTGGTAGTAAAA 3'

(SEQ ID NO: 21)

(corresponding to nucleotides 1492-1511

within exon 9 (Figure 4)

Primers 7F and 9R2 were expected to generate a fragment of ~619 bp (Figure 23) from the cDNA template. These primers will not generate a fragment from genomic DNA possibly present in the cDNA preparation, since the primers span two large introns.

mGT-3UR: 5'- GGGTTTTGGTTTTGATTGTT 3'

(SEQ ID NO: 22)

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(corresponding to nucleotides 1866-1888 within the 3' untranslated region; Figure 4).

This primer was used with primer 7F to generate the DNA fragment used in the control spike PCRs.

Primers to detect mouse ferrochelatase cDNA (EcoRI linkers, underlined):

FC-F: 5'- CTGAATTCATGTTAAACATGGGAGGCCCC 3'
(SEQ ID NO: 23)

10 (corresponding to nucleotides 215-235, Taketani et al., J. Biol.Chem. <u>265</u>: 19377-80 (1990)).

gFC-R: 5'- CTGAATTCTGCCCACTCCCTGCCGATG 3'
(SEQ ID NO: 24)

(corresponding to nucleotides 888-908, Taketani et al., J. Biol.Chem. <u>265</u>: 19377-80 (1990)).

These primers were expected to generate a 709 bp fragment (Figure 23). These primers will not generate a 20 fragment from genomic DNA possibly present in the cDNA preparation, since the primers span five introns.

Reaction volumes were 50 μ l, consisting of 4 μ l of the first strand cDNA synthesis reaction, 100 ng of each primer, 2 mM MgCl₂, 0.3 mM dNTPS, 2U of Taq-Polymerase (Bresatec) and Taq reaction buffer (Bresatec) at 1X concentration. Reactions were heated at 94°C for 2 min, then 29 cycles of 94°C for 15 sec, 58°C for 30 sec and 72°C for 1 min followed by single steps of 72°C for 4 min and 4°C for 5 min. A 10 μ l aliquot of each PCR was electrophoresed on a 2% agarose gel and DNA fragments were visualized on a UV light box after staining the gel with ethidium bromide.

Figure 24 shows the PCR fragments generated from RNA isolated from kidney (K), heart (H) and liver (L) of a

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wild-type mouse, and mice heterozygous or homozygous for the interrupted α-1,3-Gal T allele. Figure 24(i) shows that the 709 bp ferrochelatase fragment was generated from each of the cDNA preparations, indicating that cDNA template was produced from the reverse transcription reaction, and was available for the α-1,3-Gal T gene primers. The 619 bp α-1,3-Gal T fragment was present in each of the reactions spiked with the 983 bp α-1,3-Gal T cDNA product (Figure 24(ii)), indicating that amplification of the α-1,3-Gal T cDNA (spike) template had occurred.

In the reactions that were not spiked (Figure 24 (iii)), the 619 bp α -1,3-Gal T fragment was detected in cDNAs synthesized from the wild-type and heterozygous RNAs. This indicates that α -1,3-Gal T mRNA is present in the kidney, heart and liver of the wild-type and heterozygous mice. The 619 bp fragment was not detected in the unspiked homozygous KO reactions, indicating that α -1,3-Gal T mRNA is not synthesized in the homozygous KO mice.

II. TEST FOR EXPRESSION OF THE GAL EPITOPE IN HOMOZYGOUS

KNOCKOUT MICE USING ANTI-GAL ANTIBODIES WITH
FLUORESCENCE-ACTIVATED CELL SORTING (FACS)

A. Solutions

Solutions 1 to 5 are 10x isotonic.

- 1. 1.68M NaCl (948.21g/l) Dry salts overnight in
 25 hot oven before weighing
 - 2. 1.68M KCl (125g/l) Dry salts overnight in hot oven before weighing
 - 3. 1.12M CaCl_2 (165g/l $\operatorname{CaCl}_2\operatorname{2H}_2\operatorname{O}$) Dry salts overnight in hot oven before weighing
- 30 4. 1.68M $MgSO_4(414g/l MgSO_47H_2O)$ Do not dry in hot oven
 - 5. Potassium phosphate buffer pH 7.2:

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- a) 1.68M KH_2PO_4 (229 g/L)
- b) 1.12M $\rm K_2HPO_4$ (226 g/L $\rm K_2HPO_4$ $\rm 3H_2O$ or 195 g/l $\rm K_2HPO_4)$

Potassium phosphate buffer is prepared by
5 mixing together equal volumes of solutions a) and b). To
pH the buffer, remove a small sample, dilute 1:50 and
read on pH meter.

- 6. Hepes buffer 1M (CSL, Melbourne Australia)
- 7. KDS BSS:
- Add stock solutions in the following order to double-distilled water (DDW):

	Stock	Ratio of Solutions
	DDW	1210
	NaCl	121
15	KCL	3
	CaCl ₂	3
	MgSO ₄	1
	Potassium phosphate buf	fer 2
	Hepes	20

- 20 Filter sterlise, store at 4°C
 - 8. KDS/BSS/2%HSA/0.02% azide:

KDS/BSS 244.5ml
Human serum albumin 5ml
(CSL, Melbourne, Australia)
10% Na azide in MT-PBS 0.5ml

- 9. FITC dilution: Dilute 7.5ul FITC-IgG to 600ul with KDS/BSS
 - 10. Red cell lysis buffer:

0.168M NH₄C1 in double distilled water

30 11. 4% paraformaldehyde (PFA)

Solutions:

A. $NaH_2PO_42H_2O$ 22.6 g/L B. NaOH 25.2 g/L C. 40% paraformaldehyde:

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1) 4 g paraformaldehyde (BDH, Kilsyth, Australia, #29447) dissolved in 10ml double distilled water. Heat 70°C 2 hours on stirrer in fume hood and a few drops of 2M NaOH are added until the solution becomes clear.

2) 0.54 g glucose is then added.

3) Store RT in light proof bottle.

D. Add together 83 ml of A + 17 ml of B.

E. Final 4% PFA fixative solution: 90 ml of D + 10 ml of C. pH 7.4 - 7.6; adjust pH with 1M HC1.

12. Hanks Balanced Salt Solution (Ca and Mg free) (HBBS):

15 KCL 400mg KH₂PO₄ 60mg NaC1 **8**q NaHCO3 350mg Na₂HPO₄2H₂O 68mg 20 Glūcose 1g H₂O to 1 liter

adjust to pH 7.0; filter sterilize

13. Sheep antihuman IgG and IgM fluorescein
isothiocyanate (FITC) F(ab)2 fragments (Silenus,
25 Hawthorn, Australia):

B. Methods

- 1. Eye bleed mice, collect 300-400ul into prechilled Ependorf tube, store on ice, add EDTA 20mg/ml to give final concentration of 2mg/ml.
- 2. Transfer blood (including appropriate human controls) to 10ml plain tube and add 10ml red cell lysis buffer (0.168M NH₄Cl) pre-warmed to 42°C; incubate for several minutes or until cells have lysed.
- 3. Pellet cells by centrifugation (800 x g, 7 min, 35 4°C).
 - 4. Resuspend cells in 10ml KDS/BSS/2% HSA/0.02% NaN_3
 - 5. Pellet cells as above; repeat steps 4 & 5.
- 6. Resuspend cells in 1000ul KDS/BSS/2% HSA/0.1% 40 NaN_3 ; transfer aliquots to V bottom FACS tubes.

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- 7 Pellet cells as above.
- 8. Resuspend cells in 100ul KDS/BSS/2% HSA/0.1% NaN_3
- 9. Add 50ul of purified anti-GAL antibody (see 5 Example 1, above), normal human serum (NHS) or HBBS/2% HSA/0.1% NaN₃ and incubate 45 min.
 - 10. Add 2ml KDS/BSS/2% HSA/0.02% NaN3; centrifuge cells as above.
- 11. Add 50ul of a 1:80 dilution of sheep antihuman 10 IgG or IgM FITC F(ab)2 fragment (Silenus).
 - 12. Add 2ml KDS/BSS/2% HSA/0.02% NaN3; centrifuge cells as above.
 - 13. Resuspend cells in 300ul KDS/BSS/2% HSA/0.02% NaN3.
- 14. Transfer samples to plastic round-bottom FACS tubes and add 3 ul of propidium iodide (100ug/ml); samples are now ready for analysis; keep on ice.
 - 15. Analyse on Beckman FACS scan using peripheral blood lymphocyte settings.

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C. <u>Results</u>
The results of these experiments are given below:

		median channel fluorescence (log scale)	peak channel fluorescence (log scale)
5	MOUSE 129 (Normal) PBL + FITC anti- IgG alone (neg. control)	9	9
	MOUSE 19 PBL (wild type) GAL IgG	197	286
10	MOUSE 21 PBL (Gal KO) GAL IgG	22	15
15	MOUSE 129 (Normal) PBL + FITC anti- IgM alone (neg. control)	7 .	1
	MOUSE 19 PBL (wild type) GAL IgM	120	167
20	MOUSE 21 PBL (Gal KO) GAL IgM	34	18
25	MOUSE 129 PBL (normal) PBL + FITC IGG alone (neg. control)	8	9
	MOUSE 129 PBL (normal) GAL IgG	120	328
30	MOUSE 9 PBL (Gal KO) GAL IGG	10	9

The results of human anti-Gal binding to human peripheral blood lymphocytes (negative control) are not shown but were negative. These experiments demonstrate that human anti-Gal (IgG and IgM) antibodies bind to peripheral blood cells of the homozygous \$\alpha\$1,3 galactosyltransferase knockout mice (mouse 21 and mouse 9) very weakly if at all. This confirms the expected lack of the galactose \$\alpha\$1,3 galactose (GAL) epitope in

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such mice. In contrast, peripheral blood cells of normal mice (mouse 129 and mouse 19) of the same strain display clear binding of anti-Gal antibodies.

III. TEST FOR EXPRESSION OF THE GAL EPITOPE IN HOMOZYGOUS

KNOCKOUT MICE USING IB₄ LECTIN WITH FACS

 IB_4 Lectin has an exclusive affinity for terminal $\alpha-$ D-galactosyl residues, and is demonstrated below to be useful for characterizing the knockout mice.

A. Solutions

1. 4% paraformaldehyde (see above)

Mouse Tonicity PBS (MT-PBS)

Na₂HPO₄

2.27g

NaH₂PO₄2H₂O

0.62q

NaC1

8.7q

15

Make up to 1 liter with DDW

3. Dead Cell Removal Buffer (DCRB):

-4.5 g Sorbitol

-7.6 g Glucose monohydrate, (6.93 g if

anhydrous)

20

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-12.5 ml KDS/BSS

-Make up to 100 ml with DDW

-Filter, store at 4°C

-Open only under sterile conditions

- 4. KDS/BSS (Mouse Tonicity, Hepes Buffered Balanced Salt Solution pH 7.2) (see above)
- Red cell lysis buffer (see above)
- 6. KDS/BSS/2%HSA/0.02%azide (see above)
- 7. Hanks Balanced Salt Solution (Ca and Mg free) (see above)

30 B. Methods

1. Remove spleen, hold with curved forceps and collect splenocytes by injecting with a 27 gauge needle bent at 90°C, injecting (2.5 ml syringe) 100-200

ul buffer into the spleen two or three times. Using the flat surface of the bent needle massage cells out of holes made in spleen. Repeat injections and removal of cells until no cells remain in capsule.

- 5 2. Transfer splenocytes to 10ml tube and centrifuge to pellet cells (500xg, 7 min, 4°C).
- 3. Remove supernatant and add 3ml red cell lysis buffer pre-warmed to 42°C; incubate for several minutes or until cells have lysed. Underlay with 1ml 10 HIFCS (heat inactivated fetal calf serum) and stand on ice 5 minutes. Top to 10ml with KDS BSS/10% HIFCS.
 - 4. Centrifuge as above.
 - 5. Resuspend cells in 3ml dead cell removal buffer; mix well with pipette.
- 6. Pass through a glass pipette plugged with cotton wool and collect cells into a 10ml tube. Don't force cells through, allow to drain under gravity.
 - 7. Underlay cells with 1 ml BSS/10% HIFCS.
 - 8. Centrifuge as above.
- 9. Remove supernatant.
 - 10. Centrifuge as above; repeat steps 4 & 5.
 - 11. Add 0.5 ml cold 4% paraformaldehyde (PFA).
 - 12. Incubate on ice for 5 min with intermittent mixing.
- 25 13. Add 2 ml ice cold HBBS and centrifuge as above.
 - 14. Repeat washings with 2ml and then 1ml HBBS.
 - 15. Resuspend cells in 100ul KDS/BSS/2%
- 30 HSA/01.% NaN3; transfer to V bottom FACS tubes.
 - 16. Add FITC IB4 lectin (Sigma, Cat. No. L 2895), 50ul at 20ug/ml, or 50ul HBBS; incubate on ice for 30 min.
- 17. Add 2ml KDS/BSS/2% HSA/0.1% NaN₃; spin 35 cells as above.

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- 18. Resuspend cells in 300ul KDS/BSS/2% HSA/0.1% NaN3.
- 19. Transfer samples to plastic round-bottom FACS tubes; samples are now ready for analysis; keep on 5 ice.
 - 20. Analyse on FACS scanner using peripheral blood lymphocyte setting.

C. Samples

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- 1. Mouse 129 splenocytes alone
- 2. Mouse 129 splenocytes + IB4 lectin
- 3. human PBL alone
- 4. Human PBL + IB4 lectin

D. <u>Results</u>

Results of these experiments are given below:

		mean fluorescence channel (log scale)	median fluorescence channel (log scale)	peak fluorescence channel (log scale)
15	splenocytes alone (autofluorescence)	1	1	1
	mouse 19 (wild type) splenocytes	· 252	.58	16
20	mouse 21 (KO mouse) splenocytes	3	2	1

The results demonstrate that IB_4 lectin binds spleen cells of the homozygous $\alpha 1,3$ galactosyltransferase 25 gene targeted (Gal KO) mouse (mouse 21) very weakly if at all. This confirms the expected lack of the galactose $\alpha 1,3$ galactose (GAL) epitope in such mice. In contrast, peripheral blood cells of a normal mouse (mouse 19) of the same strain binds IB_4 lectin strongly.

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- IV. IMMUNOHISTOLOGICAL ASSESSMENT OF MOUSE TISSUES FOR THE PRESENCE OF THE GAL EPITOPE USING ANTI-GAL ANTIBODIES.
 - A. Reagents
 - 1. TBS: Tris Buffered Saline

NaC1

8g

KCl

0.2g

Tris base

3g

- dissolve in 800ml distilled water. Adjust pH 10 to 8.0 with 1 M HCl. Adjust volume to 1L. Sterilise by autoclaving. Store at RT.
 - 2. Blocking buffer:
 - TBS + 2% bovine serum albumin (BSA) + 10% rabbit serum:
- 15 3. Peroxidase conjugates:

DAKO (Denmark) peroxidase (POD) conjugated to rabbit anti-human IgG (fragment) and DAKO (Denmark) peroxidase (POD) conjugated to rabbit anti-human IgM (fragment).

- Conjugates were both separately pre-absorbed on 10% mouse liver powder at 4°C overnight, then centrifuged at 18,000xg for 10 minutes in a Biofuge and then at 30 psi for 30 min in a Beckman airfuge. Conjugated antisera were diluted 1/50 in 2% blocking buffer (TBS + 2% BSA + 2% rabbit serum) with 16% normal mouse serum.
 - 4. Mouse liver powder preparation:
 As modified from Antibodies, a Laboratory
 Manual Ed Harber and David Lane, Cold Spring Harbour
 Laboratories (1988) p663:
- a) Prepare a fine suspension of mouse liver in mouse tonicity phosphate buffered saline (MT-PBS).

 Mash liver through a sieve with a 5 ml plunger. Discard any fibrous tissue. One gram of tissue should be resuspended in approximately 1 ml MT-PBS.

- b) Transfer the tissue/saline suspension to ice for 5 min.
- c) Add 8 ml of acetone (-20°C) (Univar 6, Ajax Chemicals) for 10 minutes. Mix vigorously.
- 5 Incubate on ice for 30 minutes with occasional vigorous mixing.
 - d) Collect the precipitate by centrifugation at 10,000g (9,000 rpm in Sorvall RC-5B refrigerated superspeed centrifuge). Spin for 10 minutes.
- e) Resuspend the pellet with fresh acetone (-20°C) and mix vigorously. Allow to sit on ice for 10 minutes.
- f) Centrifuge at 10,000g for 10 minutes.

 Transfer the pellet to a clean piece of filter paper.

 15 Spread the precipitate and allow to air-dry at room temperature.
- g) After the pellet is dry, transfer it to an airtight container. Remove any large pieces that will not break into a fine powder. Dessicate and store at 20 4°C.

Yield as approximately 10-20% of the original wet weight. To use acetone powders, add to a final concentration of 1%. Incubate for 30 min at 4°C.

Spin at 10,000g for 10 minutes. (13,000 rpm in Biofuge)

25 5. DAB/H₂O₂/Imidazole:

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Peroxidase substrate: 3,3'-Diaminobenzidine tetrahydrochloride (DAB) (Sigma, Missouri)

- 1 tablet DAB (take out of fridge 10 min before use)
- 1 tablet urea H₂O₂ (Sigma, Missouri)
 - add to 15 ml tris HCL, pH 7.6 + 0.01M imidazole (0.0102g), (Sigma, Missouri)
 - make up immediately before use
 - 6. Tris HCL:

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1.211g Tris in 200ml double distilled water pH 7.6

7. Animal serum sources:

Mouse and rabbit sera were obtained in-house

5 (St. Vincent's Hospital, Dept, of Clinical Immunology).

Sheep serum was obtained from the University of
Melbourne Veterinary Clinic and Hospital, Werribee,
Australia.

8. Harris Haematoxylin:

Haematoxylin C.I. 75290 (BDII, Poole, U.K.

#34037) 10g

Absolute ethanol 200ml
Potassium alum 200g
double distilled water 2000ml

15 glacial acetic acid 80ml

Preparation:

1. Dissolve

haematoxylin in absolute ethanol

2. Heat to dissolve alum in double distilled water

3. Mix solution 1 and 2

4. Immediately before use add 80 ml 1% sodium iodate and 80 ml glacial acetic acid

9. Scott's Tap Water:

Sodium hydrogen carbonate 14 g
MgSO₄ 80 g

Tap water

4 litres

B. Methods

- 1. Cut 4 um sections of the relevant tissue on cryostat
- 30 2. Tissue should be free of cracks
 - 3. Air dry slides for 30 min

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- 4. Apply 10% blocking buffer at room temp in humidified chamber, 60 min
- 5. Remove blocking antibody with tissue made to fine point
- 6. Apply 1st antibody, anti-GAL, or 2% blocking buffer as control, 50ul, ensure no air bubbles and incubate at room temp in humidified chamber for 30 min
 - 7. Wash off with Tris buffered saline (TBS) 3 times 2 minutes washes
 - 8. Apply second antibody 1:50 peroxidase (POD) conjugated rabbit anti-human IgG and IgM (DAKO, Denmark); incubate 30 min at room temp in humidified chamber
- 9. Wash off with Tris buffered saline (TBS) 3 times 3 minute washes
 - 10. Wash off with TBS as above
 - 11. Incubate DAB/H₂O₂/imidazole for 10 minutes
 - 12. Wash in water
- 20 13. Stain with haemotoxylin C 10 seconds
 - 14. Wash in water
 - 15. Place in Scotts tap water for 15 seconds
 - 16. Wash in water
 - 17. Wash in absolute alcohol (x3) (Univar 214, Ajax chemicals)
 - 18. Wash in absolute xylene (x3) (Univar 577, Ajax chemicals)
 - 19. Coverslip using automatic coverslip machine (Tissue Tek)

30 Controls:

Buffer only + POD conjugated rabbit anti-human IgM (negative)

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- 2. Buffer only + POD conjugated rabbit anti-human IgG (negative)
- 3. Human kidney (negative)
- 4. Pig renal cortex (positive)

5 Samples:

- 1. Mouse 129 SV (control) kidney
- 2. mouse 9 (Gal Knockout) kidney
- 3. mouse 21 (Gal Knockout) kidney

C. Results

10 KIDNEY

	GLOMERULI	ENDOTHELIUM	comments
MOUSE 129 anti-IgM	POSITIVE	POSITIVE	
MOUSE 9 anti-IgM	NEGATIVE	NEGATIVE	weak adventitial staining
MOUSE 21 anti-IgM	NEGATIVE	NEGATIVE	weak adventitial staining
MOUSE 129 anti-IgG	POSITIVE	POSITIVE	7
MOUSE 9 anti-IgG	NEGATIVE	NEGATIVE	
MOUSE 21 anti-IgG	NEGATIVE	NEGATIVE	·
POD conjugated antibody alone	ALL NEGATIVE	ALL NEGATIVE	

These results indicate that human anti-Gal IgG and IgM antibodies do not bind kidney tissue of the α1,3 galactosyltransferase gene targeted (Gal KO) mice (mouse 21 and mouse 9). This confirms that lack of the galactose α1,3 galactose (GAL) epitope in the gene targeted (KO) mice. In contrast, these antibodies react strongly with the endothelium of blood vessels and the glomeruli of a normal mouse of the same strain (129).

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- V. IMMUNOHISTOLOGICAL EXAMINATION OF MOUSE TISSUES USING IB_4 LECTIN
 - A. Reagents

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- 1. Blocking buffer: TBS + 2% BSA + 10% sheep serum
- 2. FITC IB₄ (Sigma, Missouri, USA #L-2895)

1 mg diluted in 1 ml HBBS to give stock solution, then dilute to final volume of 20 ug/ml in TBS + 2% BSA + 2% sheep serum

- 3. Peroxidase anti-FITC
- Boehringer anti-fluorescein POD Fab fragments; dilute 1/300 in 2% blocking buffer
 - 4. DAB/H₂O₂/Imidazole see above
 - 5. Tris HCL see above
 - 6. Animal serum sources see above
- 7. Harris Haematoxylin see above
 - 8. Scott's Tap Water see above
 - B. Methods
 - Preparation of Sections; same as Section 4B, steps 1-7 above.
- 20 2. Apply 50 μ l FITC conjugated IB4 (Sigma # 1-2894) 20 μ g/ml, incubate in a humidified chamber for 30 minutes.
 - Wash with TBS, 3 minutes (x3).
- 4. Apply 50 μ l per oxidase conjugated anti FITC Fab fragments (Boehringer Mannheim), diluted 1-3-- with TBS + 2% BSA + 2% sheep serum. Incubate for 30 minutes in humidified chamber.
 - 5. Wash with TBS, 3 minutes (x3).
- 6. Processing for microscopy same as Section IVB steps 14-22.

Controls

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1.	Buffer only + POD anti-FITC	(negative)
2.	Human kidney	(negative)
3.	Pig renal cortex	(positive)

Samples 1st Experiment

2	Mousemouse	6	wild type	heart	liver	kidney kidney	Juna
	mousemouse		neterozygote KO	heart	liver	kidney kidney	lung
		_	monday wo	mear c	TIVEL	Klanev	inna

Samples 2nd Experiment

10			wild type heart liver kidney lung
	2.	mouse 20	heterozygote KO heart liver kidney lung
	З.	mouse 21	homozygous KO heart liver kidney lung

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C. Results

Kidney

	GLOMERULI	ENDOTHELIUM
HUMAN	NEGATIVE	NEGATIVE
PIG	POSITIVE	POSITIVE
129 MOUSE	POSITIVE	POSITIVE
MOUSE 6	POSITIVE	POSITIVE
MOUSE 7	POSITIVE	POSITIVE
MOUSE 9	NEGATIVE	NEGATIVE
MOUSE 19	POSITIVE	POSITIVE
MOUSE 20	POSITIVE	POSITIVE
MOUSE 21	NEGATIVE	NEGATIVE
anti-FITC alone	ALL NEGATIVE	ALL NEGATIVE

Liver

	ENDOTHELIUM	BILE DUCT
129 MOUSE	POSITIVE	POSITIVE
MOUSE 6	POSITIVE	POSITIVE
MOUSE 7	POSITIVE	POSITIVE
MOUSE 9	NEGATIVE	NEGATIVE
MOUSE 19	POSITIVE	POSITIVE
MOUSE 20	POSITIVE	POSITIVE
MOUSE 21	NEGATIVE	NEGATIVE
anti-FITC alone	ALL NEGATIVE	ALL NEGATIVE

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Heart

	ENDOTHELIUM	PERINUCLEAR	ENDO- MYOCARDIUM
129 MOUSE	POSITIVE	POSITIVE	POSITIVE
MOUSE 6	POSITIVE	POSITIVE	POSITIVE
MOUSE 7	POSITIVE	POSITIVE	POSITIVE
MOUSE 9	NEGATIVE	NEGATIVE	NEGATIVE
MOUSE 19	POSITIVE	POSITIVE	POSITIVE
MOUSE 20	POSITIVE	POSITIVE	POSITIVE
MOUSE 21	NEGATIVE	NEGATIVE	NEGATIVE
anti-FITC alone	ALL NEGATIVE	ALL NEGATIVE	ALL NEGATIVE

10 Lung

	ENDOTHELIUM	BRONCHI	PARENCHYMA
129 MOUSE	POSITIVE	POSITIVE	POSITIVE
MOUSE 6	POSITIVE	POSITIVE	POSITIVE
MOUSE 7	POSITIVE	POSITIVE	POSITIVE
MOUSE 9	NEGATIVE	NEGATIVE	NEGATIVE
MOUSE 19	POSITIVE	POSITIVE	POSITIVE
MOUSE 20	POSITIVE	POSITIVE	POSITIVE
MOUSE 21	NEGATIVE	NEGATIVE	NEGATIVE
anti-FITC alone	ALL NEGATIVE	ALL NEGATIVE	ALL NEGATIVE

These results indicate that IB₄ lectin does not bind kidney, heart, liver or lung tissue of the α1,3 galactosyltransferase gene targeted (Gal KO) homozygous mice (mouse 21 and mouse 9). This confirms the lack of the galactose α1,3 galactose (GAL) epitope in the gene targeted (KO) mice. In contrast these antibodies react strongly with the tissues of a normal mouse and heterozygous KO mice (mouse 129, mouse 6, mouse 7, mouse 19, mouse 20) of the same strain.

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VI. RESISTANCE OF SPLEEN CELLS FROM KNOCKOUT MICE TO LYSIS BY HUMAN SERUM

Lysis of spleen cells by human serum was tested through use of a 51 chromium release assay. See in general Example 4, above.

- A. Preparation of Mouse Splenocytes Shortman, K.J. et al, Immunological Methods. 1:273-287 (1972).: -Dissect out spleen, avoid damaging outer membranes and carefully remove mesentery tissue and fat.
- -Place in petri dish, with 1 ml RPMI 1640 (Gibco BRL)
 /10% Heat-inactivated foetal calf serum (HI-FCS). (Heat-inactivation = 40 Min at 56°C).
 -Gently tease out cells into petri dish, collect and centrifuge 500xg, 5 min, 4°C
- -Remove RPMI/10% HIFCS, gently resuspend cells in 3 ml 0.9% NH4C1 (0.168M), using a Pasteur pipette. (Use Pasteur pipettes or wide-bore pipettes for all re suspension and transfer procedures)
- -Transfer to 10 ml tube, underlay with 1 ml HIFCS, stand on ice, 5 min.
 - -Transfer supernatant to clean tube, centrifuge 500xg, 7 min, 4°C
 - -Discard supernatant, re-suspend cells in 3 ml dead cell removal buffer, mix well with pipette.
- -Pass through cotton wool plug in glass pipette (under gravity, do not force through), collect cells into 10 ml tube.
 - -Underlay cells with 1 ml HI-FCS.
 - -Centrifuge 500xg, 7 min, 4°C
- -Remove supernatant, re-suspend cells in 50 μ l RPMI, 10% HI-FCS. Store cells on ice.
 - B. Preparation of Serum:
- Human Collect whole blood from a pool of normal donors; allow to stand at room temp. for 2 hours.

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Wring the clot with an 'Orange stick'; spin Remove and pool serum. Store half at -70°C in 3 ml aliquot's (normal human serum); heatinactivate the other half, see below.

5 Fetal calf serum - purchased from Gibco BRL, and stored at -20°C.

della di

- C. Cell Counting:
- 1. Add 5 μ l cells to 95.0 μ l RPMI, 10% HI-FCS
- 2. Remove 10 μ l cells, add 10 μ l Acridine
- Orange/Et Br solution, (Lee, S.K. et al. Eur J. Immunol. 1975. 5: 259-262)
 - Count cells, (viable = green, deads = orange).
 - 4. Cell viability should be approx. 90-100 %
 - 5. Calculate cell number.
- D. 51Chromium Labelling:

Cell Type

; .

Incubation conditions

Time Amount⁵¹Cr/10⁷ cells

Freshly prepared cells: ~2 hours ~150-300 μ C1 (eg., splenocytes or

20 lymphocytes)

Cultured Cells:

~1 hour $\sim 100 \,\mu\text{C}1$

Labelling:

Combine:

- cells (2 X 10⁷)
- (⁵¹Cr) Sodium Chromate in 0.9% NaCl solution (the volume added depends on cell type as indicated above and on the specific activity of the - (⁵¹Cr) Sodium Chromate).

- RPMI/2% HIFCS up to a total of 200 μl

Incubate at 37°C for time shown above with gentle agitation every 15 min.

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E. Washing

- -Place 4ml HI-FCS into 10ml tube and carefully layer labelling reaction on top with a swirling motion; centrifuge 5 min, 500xg, 4°C.
- 5 -Remove top two layers with a careful circular motion using a glass pipette.
 - -Resuspend cells in 1ml RPMI/2% HI FCS
 - -Pellet cell suspension through another 4 ml HI FCS
- -Resuspend cell pellet in 1ml RPMI/2% HI FCS, store on ice.

F. Release Assay:

-Perform assay in 96-well microtire plate (ICN-FLOW).

-Assay should be set up in quadruplicate.

15 -Assay is performed in a total volume of 180 μ l. Assay:

		NHS	*HI-NHS	16% SDS	CELLS	RPMI/2% HIFCS
	MAX Release	_	90µ1	22.5µl	$25\mu 1$	42.5µl
	Spont.Release	_	90		25	65µ1
20	5% NHS	9μ l	81		25	65
	10% NHS	18	72		25	65
	20% NHS	36	54		25	65
	30% NHS	54	36		25	65
	40% NHS	72	18		25	65
25	50% NHS	90	_		25	65

- *HI = heat inactivated
- -All volumes indicated are in μ l
- -Reaction components are added to the plate in the order: RPMI, Serum and ⁵¹Cr-labelled cells.
- 30 -Cover plate with plate-sealer
 - -Incubate, 4 hours, 37°C.
 - -Spin plate, 1500 rpm, 5 min.
 - -Remove plate-scaler, remove 80 μ l from each wall, count released chromium on gamma counter.
- -Calculate specific lysis for each well according to the formula:
 - % Specific Lysis = (Test cpm Spontaneous release cpm) X 100
 (Maximal release cpm Spontaneous release cpm)

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Calculate mean and standard deviation for each experimental point. Graph % Human serum (X axis) against % Specific lysis (Y axis) for each type of cell (wild type, heterozygote KO and homozygous KO)

The results of these experiments are depicted in Figure 25. The results indicate that spleen cells from a homozygous knockout mouse are relatively resistant to lysis by human serum, in comparison to spleen cells derived from mice heterozygous for the interrupted allele or from wild-type mice.

EXAMPLE 16

Generation of Knockout Animals Through Microinjection of Eqqs

Transgenic animals are generated routinely 15 by microinjection of DNA into the pronuclei of fertilised eggs. Generally this technology results in the random integration of the transgene in the genome. conventional transgenic technology has resulted in homologous recombination between the injected transgene 20 and the endogenous gene. See, for example, Brinster et al., Proc. Nat. Acad. Sci. USA 86: 1087-91 (1989). Described below are procedures for inactivating the α -1,3-Gal T gene in pigs through microinjection of eggs with gene targeting constructs.

25 I. GENE TARGETING CONSTRUCTS

The frequency of homologous recombination in embryos is improved if the gene targeting constructs are prepared with isogenic DNA. Therefore the "knock out" constructs are prepared from DNA isolated from the boar used to fertilize the oocytes used for microinjection. DNA is isolated from the tail or ear tissue, and genomic fragments from both $\alpha-1,3$ -Gal T alleles of the boar, encompassing exons 8 & 9 are cloned using long range PCR or conventional genomic library 35 technologies. Clones for each of the $\alpha-1,3$ -Gal T alleles

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are identified using restriction fragment length polymorphism identification and DNA sequencing. Constructs to target both alleles are made by interrupting the coding sequence of exon 9, either by deletion or by inserting a heterologous DNA fragment. The constructs contain at least 8 kb of homologous DNA to promote efficient homologous recombination.

Various approaches can be used to detect gene targeting events, depending on the strategies used in designing the knockout constructs. Several such approaches, and the corresponding strategies for construction of constructs, are provided below:

a) PCR of Genomic DNA:

Homologous DNA on one side of the interrupting DNA fragment is constructed to be less than 1 kb, allowing PCR amplification of a short diagnostic fragment. (Amplification of small fragments generally is relatively efficient).

b) Reverse Transcription/PCR:

A deletion of about 100 bp within exon 9 is made, allowing synthesis of a shortened α-1,3-Gal T mRNA in correctly targeted cells. The shortened mRNA is detected by RT/PCR, using primers that amplify a fragment extending from exon 8 and encompassing the deletion site.

c) Green Fluorescent Protein (GFP) gene _expression:

GFP is a protein from the bioluminescent jelly fish Aequorea victoria. It absorbs blue light (395 nm) and fluoresces to emit green light (509 nm). GFP is a useful marker for gene expression. Chafie et al., Green Fluorescent Protein as a Marker for Gene Expression. Science 263: 802-5 (1994). The α -1,3-Gal T gene is interrupted within exon 9 by in-frame insertion of the GFP coding region. Expression of the GFP gene (with

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resulting fluorescence at 509 nm) is driven by the α -1,3-Gal T gene promoter in correctly targeted cells.

II. GENERATING EMBRYOS FOR MICROINJECTION

Fertilized embryos are generated as described by Nottle et al., (1993). Proc Aust Soc for Reproductive Biol 26, 33. The protocol involves:

a) Sperm from the boar providing DNA for the targeting construct is collected and stored frozen in liquid N_2 .

10 b) Superovulation of donor gilts:

Gilts are mated at the second oestrus, and aborted between days 25-40 days of gestation to synchronise the subsequent oestrus cycles. Abortion is achieved by intramuscular injection of 1 mg cloprostenol (a prostaglandin $F2\alpha$ analogue), followed by a second 0.5 mg injection 24 hours later. Gilts are superovulated by injection of 1000 i.u. equine chorionic gonadotrophin (eCG) or pregnant mare serum gonadotrophin at the time of the second cloprostenol injection, and a subsequent injection 72 hours later of 500 i.u. human chorionic gonadotrophin (hCG).

c) Fertilization:

Superovulated gilts are artificially inseminated 20-30 hours after the hCG injection, followed by a second insemination 2-4 hours later, with semen from the boar that provided DNA for the targeting construct.

d) Embryo collection:

Embryos are collected surgically 50-56 hours after hCG injection prior to fusion of the pronuclei. Oviducts are flushed with 15-20 ml phosphate saline buffer containing 1% fetal calf serum. One-cell embryos are recovered by searching oviductal flushings using low magnification microscopy.

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III. MICROINJECTION OF EMBRYOS

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Embryos are centrifuged at 12000 x g for 8 min to stratify the cytoplasm and allow the pronuclei to be visualised, and held in Dulbecco's Minimal Essential Medium with 25 mM Hepes and 5 mg/ml bovine serum albumin. Pronuclei are injected, using differential interference contrast optics, with 4-10 picolitres of DNA (10 ng/ μ l) in PBS. Gene targeting with isogenic DNA is maximized by coinjecting both allelic constructs derived from the boar into the male pronucleus.

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IV. TRANSFER OF INJECTED EMBRYOS TO RECIPIENT GILTS

The oestrus cycles of recipient gilts are synchronized with those of donors. The recipients are mated and aborted using the protocol described above, and injected with 500 i.u. eCG. Injected embryos are transferred surgically (20-40 per oviduct) to recipients on the same day that they are collected from donor gilts.

V. SCREENING FOR HOMOLOGOUS RECOMBINATION

Homologous recombinants can be detected by analysis of tissue from the born piglets. Screening procedures involve PCR technology, the precise strategy depending on the design of the gene targeting construct. Because many α-1,3-Gal T mRNA molecules are synthesized from a single α-1,3-Gal T gene in expressing cells, the RT/PCR approach can be more sensitive than PCR amplification of genomic DNA. The RT/PCR screening strategy relies on successful transcription of the interrupted gene and relative stability of the shortened mRNA.

Alternatively, constructs that promote expression of heterologous genes (eg: GFP) in correctly targeted cells allow embryos to be screened at the blastocyst stage for marker gene expression (i.e.: GFP expression can be detected by measuring fluorescence within blastocysts at 509 nm). The microinjected embryos are cultured in vitro until blastocyst development, screened for fluorescence, and fluorescing embryos transferred into recipients.

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EXAMPLE 17

A Novel Variant of Leukemia Inhibitory Factor (LIF)

Previous reports have demonstrated the existence of two forms of murine LIF. The original form (from the D transcript) was expressed and commercialized by AMRAD Corporation Ltd (Kew Victoria, Australia). The protein product derived from this transcript (hereinafter "D-LIF") is sold commercially by AMRAD as "ESGROW". Another form of LIF (hereinafter "M-LIF"), derived from an alternative transcript, is described in US Patent Application No. 07/994,099 and in Rathjen et al., Cell 62: 1105-14 (1990). The present inventors have now found a third transcript of LIF (hereinafter "T-LIF") which is found in ES cells and in human teratocarcinoma-derived cell lines such as the GCT 27 teratocarcinoma-derived cell lines described by Pera et al., Differentiation 42: 10 (1989).

The T-LIF protein is found intracellularly in contrast to the other two forms of LIF which are both extracellular. The transcript was cloned using the RACE PCR technique (see below) from murine ES cells and human GCT 27 teratocarcinoma-derived cell lines, and sequenced using standard methods. presence of the T-LIF transcript was confirmed by PCR analysis of ES cell mRNA and RNA' ase protection on GCT 27 The transcript comprises a novel first exon, located in the first intron of the LIF gene, spliced to the known exon 2 and exon 3 sequences. The mouse nucleotide sequence (SEQ ID NO: 25) and deduced amino acid sequence (SEQ ID NO: 26) are set out in Figure 26. The human nucleotide sequence (SEQ ID NO: 31) and deduced amino acid sequence (SEQ ID NO: 32) are set out in Figure 27.

When expressed in a COS cell expression system, the murine T-LIF transcript produces a 17 kD

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protein that is unglycosylated (D-LIF is glycosylated in the Golgi during the secretion process) (Figure 28). Translation of T-LIF initiates at the first in-frame initiation codon (ATG) in exon 2 to produce a protein of 158 amino acids. The protein is 45 amino acids shorter than the unprocessed D-LIF protein and 22 amino acids shorter than the mature D-LIF product generated by cleavage of the signal sequence. Because the T-LIF protein does not contain a signal sequence, it does not leave the cell and is unglycosylated. The T form of LIF is efficacious in preventing the differentiation of ES cells in culture.

METHODS

RACE CDNA CLONING

15 Cytoplasmic RNA (10 μ g) from CP1 murine ES cells (Bradley et al., Nature 309: 255-56 (1984) was reverse transcribed from the oligonucleotide 5'ACACGGTACTTGTTGCA-3' (SEQ ID NO: 27), which hybridizes to residues 500-484 of the murine LIF cDNA. The RNA was 20 added to 20 pmol of primer and $2\mu 1$ of 10x annealing buffer (500mM Tris-HCl (pH 8.0), 60mM MgCl2, 400mM KCl) in a total volume of 16μ l, heated to 85°C for 5 min, and cooled slowly to room temperature. The elongation reaction was carried out as described by Frohman et al. 25 (Proc. Natl. Acad. Sci. USA 85: 8998-9002 (1988)). Excess oligonucleotide was removed by gel filtration through a 2ml Sephacryl S-400 (Pharmacia) column equilibrated with 0.05 x TE (TE = 10mM Tris-HCl pH 7.6, 1.0mM EDTA). Fractions of 50µl corresponding to the cDNA 30 radioactive peak were pooled, concentrated by vacuum centrifugation, and resuspended in $23\mu l$ of H_20 . the 3'-end of the cDNA with dG residues, 3μ l of 10mM dGTP and 6μ l of 5 x tailing buffer (Bethesda Research Laboratories) were added and the mixture was incubated at

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37°C for 60 min. and then at 70°C for 15 min. After ethanol precipitation, the cDNA template was resuspended in $500\mu l$ H₂0.

PCR was carried out using a mouse LIF specific oligonucleotide, 5'-TTCTGGTCCCGGGTGATATTGGTCA-3' (residues 389-365) (SEQ ID NO: 28), and an anchor oligonucleotide, 5'-CCATGGCCTCGAGGGCCCCCCCCCCC-3' (SEQ ID NO: 29). PCR was carried out in a final volume of $50\mu l$ containing $7\mu l$ of the cDNA template and 34pmol of each oligonucleotide. Reaction conditions were as recommended by Perkin-Elmer Cetus, with a final concentration of 1.5mM MgCl2. DNA was denatured prior to the addition of Taq polymerase (Perkin-Elmer Cetus) by heating the reaction mixture to 94°C for 5 min. cycle (35 in total) consisted of denaturation for 2 min at 94°C, annealing for 2 min at 55°C, and elongation for 3 min at 72°C. After the final elongation (30 min at 72°C), samples were ethanol precipitated, digested with SmaI and XhoI and analyzed by agarose gel electrophoresis. DNA was purified from agarose gels using Geneclean and cloned into Sall- and Smal- digested TST7 19U (Stratagene). Suitable recombinant plasmids were purified by the rapid boiling method.

Double-stranded sequencing was performed with Sequenase version 2.0 (USB) according to the manufacturers recommendations.

BIOLOGICAL ASSAY FOR LIF ACTIVITY

An undifferentiated, murine ES cell culture (MBL5; Pease et al., Dev. Biol. 141: 344-52 (1990), between passages 15 and 30) is trypsinized and made into a single cell suspension. The cells are pelleted by centrifugation and resuspended in complete ES Cell Medium without LIF (DMEM (without Hepes), 10% FCS, 1mM β ME, 1mM glutamine). The cells are then seeded into

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24-well microtitre plates at 5×10^2 cells/16 mm well containing 1 ml of ES Cell Medium without LIF.

The complete T-LIF open reading frame was reconstructed from the PCR product and inserted into the COS cell expression vector pXMT2 as described by Rathjen et al., Cell 62: 1105-14 (1990). The plasmid used for transfection of COS cells is shown in Figure 29. The COS cells were transfected by electroporation. Supernatants from COS cells expressing T-LIF were added to the above ES cells in various dilutions (1/5, 1/10, 1/50, 1/100, 1/50, 1/1000) and incubated for 4 days in an incubator with 10% CO₂. Controls used supernatants from COS cells expressing D-LIF (pDR1, Rathjen et al., Cell 62: 1105-14 (1990)).

LIF activity is assessed as present if cells morphologically resemble ES-cells after 4 days and are distinct from the controls incubated without any form of LIF. The ES-cells are also stained for alkaline phosphatase as undifferentiated ES-cells are positive for this marker.

Even though T-LIF is produced intracellularly, sufficient numbers of cells lyse to give significant amounts of LIF activity in the culture supernatants. If the COS cells expressing T-LIF are lysed, more LIF activity is released.

PCR DETECTION OF T-LIF TRANSCRIPT

PCR was carried out on ES cell cDNA (prepared as described above except that the cDNA was not tailed with dG). PCR conditions were as described above except that 2mM MgCl₂ was used in the reactions. The oligonucleotides 5'-CACCTTTCGCTTTCCT-3' (SEQ. ID NO. 30) and 5'-TTCTGGTCCCGGGTGATATTGGTCA-3' (SEQ. ID. NO 28) were used at 80 picograms/reaction. Products of the PCR reaction were ethanol precipitated as described above,

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separated electrophoretically on a 2% agarose gel and transferred to a nylon membrane for detection using Southern hybridization (Figure 30). The probe was the full length D-LIF transcript isolated from pDR1 (Rathjen et al., Cell 62: 1105-14 (1990). The control experiment is designed to detect all LIF transcripts using internal primers 5'-TTCTGGTCCCGGGTGATATTGGTCA-3' (SEQ. ID. NO 28) and 5'-CTGTTGGTTCTGCACTGGA-3' (SEQ. ID. NO. 33).

The foregoing detailed description has been provided for a better understanding of the invention only and no unnecessary limitation should be understood therefrom as some modifications will be apparent to those skilled in the art without deviating from the spirit and scope of the appended claims.

What is claimed is:

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- A purified and isolated nucleic acid molecule comprising a nucleic acid sequence selected from the group consisting of (1) the porcine nucleic acid sequence depicted in Figure 4 (SEQ ID NO: 7), (2) a
 sequence corresponding to the sequence of (1) within the scope of the degeneracy of the genetic code, (3) a sequence that encodes a porcine polypeptide having α-1,3 galactosyltransferase activity and that hybridizes under standard high stringency conditions with a sequence
 complementary to the sequence of (1) or (2), and (4) a sequence complementary to the sequence of (1), (2) or (3).
 - 2. A host cell that is transformed with the nucleic acid molecule of claim 1.
- 15 3. A porcine α -1,3 galactosyltransferase encoded by the nucleic acid molecule of claim 2.
- 4. A DNA construct useful for inactivating the porcine α-1,3 galactosyltransferase gene by insertion of a desired DNA sequence into an insertion site of said gene, comprising said desired DNA sequence flanked by first and second homology sequences, said first and second homology sequences, respectively, sufficiently homologous to first and second genomic sequences flanking said insertion site to allow for homologous recombination of said DNA construct with said porcine α-1,3 galactosyltransferase gene when said DNA construct is introduced into a porcine cell having said α-1,3 galactosyltransferase gene.

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- 5. The DNA construct of claim 4, wherein said insertion site is within exon 4, exon 7, exon 8 or exon 9 of the porcine α -1,3 galactosyltransferase gene.
- 6. The DNA construct of claim 4, wherein said desired DNA sequence is selected from the group consisting of the neo^R gene, the hyg^R gene and the thymidine kinase gene.
- 7. The DNA construct of claim 6, wherein said desired DNA sequence is bordered at the 5' and 3' ends by FRT DNA elements, and wherein stop codons for each of the three reading frames have been inserted 3' to the desired DNA sequence.
- 8. A DNA construct useful for inactivating the murine α-1,3 galactosyltransferase gene by insertion of a desired DNA sequence into an insertion site of said gene, comprising said desired DNA sequence flanked by first and second homology sequences, said first and second homology sequences being, respectively, sufficiently homologous to first and second genomic
 20 sequences flanking said insertion site to allow for homologous recombination of said DNA construct with said murine α-1,3 galactosyltransferase gene when said DNA construct is introduced into a murine cell having said α-1,3 galactosyltransferase gene.
- 25 9. The DNA construct of claim 8, wherein said insertion site is within exon 4, exon 7, exon 8 or exon 9 of the murine α -1,3 galactosyltransferase gene.
 - 10. The DNA construct of claim 8, wherein said desired DNA sequence is selected from the group

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consisting of the neo^{R} gene, the hyg^{R} gene and the thymidine kinase gene.

- 11. The DNA construct of claim 10, wherein said desired DNA sequence is bordered at the 5' and 3' ends by FRT DNA elements, and wherein stop codons for each of the three reading frames have been inserted 3' to the desired DNA sequence.
- 12. A method for generating a mammalian totipotent cell having at least one inactivated α-1,3
 10 galactosyltransferase allele, said totipotent cell derived from a mammalian species having a functional α-1,3 galactosyltransferase gene, comprising:
 - (a) providing a plurality of cells characterized as totipotent cells of said mammalian species;
 - (b) introducing into said totipotent cells a nucleic acid construct effective for inactivating said α -1,3 galactosyltransferase gene by insertion of a desired DNA sequence into an insertion site of said gene through homologous recombination; and
 - (c) identifying a totipotent cell having at least one inactivated $\alpha\text{--}1,3$ galactosyltransferase allele.
- The method of claim 12 in which said totipotent cell is a murine ES cell.
 - 14. The method of claim 12 in which said totipotent cell is a murine egg.
 - 15. The method of claim 12 in which said totipotent cell is a porcine ES cell.

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- 16. The method of claim 12 in which said totipotent cell is a porcine PGC.
- 17. The method of claim 12 in which said totipotent cell is a porcine egg.
- 18. A method for generating a mammal lacking a functional α -1,3 galactosyltransferase gene, said mammal belonging to a species having a functional α -1,3 galactosyltransferase gene, comprising:
- (a) providing a mammalian totipotent cell having at least one inactivated $\alpha-1$, 3 galactosyltransferase allele, said totipotent cell derived from a mammalian species having a functional $\alpha-1$, 3 galactosyltransferase gene;
- (b) manipulating said totipotent cell such that mitotic descendants of said cell constitute all or part of a developing embryo;
 - (c) recovering a neonate derived from said embryo; and
- (d) raising and breeding said neonate to obtain a mammal homozygous for said inactivated α -1,3 galactosyltransferase allele.
- 19. The method of claim 18, wherein said totipotent cell is a murine ES cell and said manipulating comprises injecting said ES cell into the blastocyst cavity of a murine blastocyst and implanting said injected blastocyst into a murine recipient female.
- 20. The method of claim 18, wherein said totipotent cell is a murine egg, and said manipulating comprises implanting said egg into a murine recipient female.

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- The method of claim 18, wherein said totipotent cell is a porcine ES cell and said manipulating comprises injecting said ES cell into the blastocyst cavity of a porcine blastocyst and implanting said injected blastocyst into a porcine recipient female.
- The method of claim 18, wherein said totipotent cell is a porcine ES cell and said manipulating comprises injecting said ES cell into a porcine morula.
- The method of claim 18, wherein said totipotent cell is a porcine ES cell and said manipulating comprises co-culture of said ES cell with a zona pellucida-disrupted porcine morula.
- The method of claim 18, wherein said totipotent cell is a porcine ES cell and said manipulating comprises fusing said ES cell with an enucleated porcine zygote.
- The method of claim 18, wherein said totipotent cell is a porcine egg, and said manipulating comprises implanting said egg into a porcine recipient female.
- 26. A mammal lacking a functional α-1,3 galactosyltransferase gene, said mammal belonging to a species having a functional α-1,3 galactosyltransferase
 25 gene, said mammal produced by the method of claim 18.
 - 27. The mammal of claim 26, wherein said mammal is a mouse.

- 28. The mammal of claim 26, wherein said mammal is a pig.
- 29. A non-naturally occurring mammal lacking a functional α -1,3 galactosyltransferase gene, said mammal belonging to a species having a functional α -1,3 galactosyltransferase gene.
 - 30. The mammal of claim 29, wherein said mammal is a mouse.
- The mammal of claim 29, wherein said mammal is a pig.
- 32. A purified and isolated nucleic acid molecule comprising a nucleic acid sequence selected from the group consisting of (1) the nucleic acid sequence depicted in Figure 26 (SEQ ID NO: 25), (2) a sequence corresponding to the sequence of (1) within the scope of the degeneracy of the genetic code, (3) a sequence that encodes murine T-LIF and that hybridizes under standard high stringency conditions with a sequence complementary to the sequence of (1) or (2), and (4) a sequence complementary to the sequence of (1), (2) or (3).
 - 33. A host cell that is transformed with the nucleic acid molecule of claim 32.
 - 34. A murine T-LIF polypeptide encoded by the nucleic acid molecule of claim 32.
- 25 35. A purified and isolated nucleic acid molecule comprising a nucleic acid sequence selected from the group consisting of (1) the nucleic acid sequence depicted in Figure 27 (SEQ ID NO: 31), (2) a sequence

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corresponding to the sequence of (1) within the scope of the degeneracy of the genetic code, (3) a sequence that encodes human T-LIF and that hybridizes under standard high stringency conditions with a sequence complementary to the sequence of (1) or (2), and (4) a sequence complementary to the sequence of (1), (2) or (3).

- 36. A host cell that is transformed with the nucleic acid molecule of claim 35.
- 37. A human T-LIF polypeptide encoded by the nucleic acid molecule of claim 35.
- 38. A method for eliminating or reducing hyperacute rejection of non-primate mammalian cells, tissues and organs by human serum, comprising adding, to said human serum, a physiologically acceptable amount of galactose or a saccharide in which the terminal carbohydrate is an α galactose linked at position 1, prior to exposure of said human serum to said non-primate cells, wherein said amount of galactose or saccharide is sufficient to reduce or eliminate said hyperacute rejection.
 - 39. The method of claim 38, wherein said saccharide is selected from the group consisting of melibiose, galactose α 1-3 galactose and stachyose.
- 40. A method for eliminating or reducing
 25 hyperacute rejection of non-primate mammalian cells,
 tissues and organs by human serum, comprising
 substantially depleting said serum of immunoglobulin.
 - 41. A method for eliminating or reducing hyperacute rejection of non-primate mammalian cells,

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tissues and organs by human serum, comprising substantially depleting said serum of IgM antibodies.

A method for eliminating or reducing hyperacute rejection of non-primate mammalian cells by human serum, comprising substantially depleting said serum of anti-GAL IgM and IgG antibodies.

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- 43. A method for eliminating or reducing hyperacute rejection of non-primate mammalian cells by human serum, comprising substantially depleting said serum of anti-GAL IgM antibodies.
- 44. Affinity-treated human serum substantially free of anti-GAL antibodies.
- 45. Affinity-treated human serum substantially free of anti-GAL IgM antibodies.

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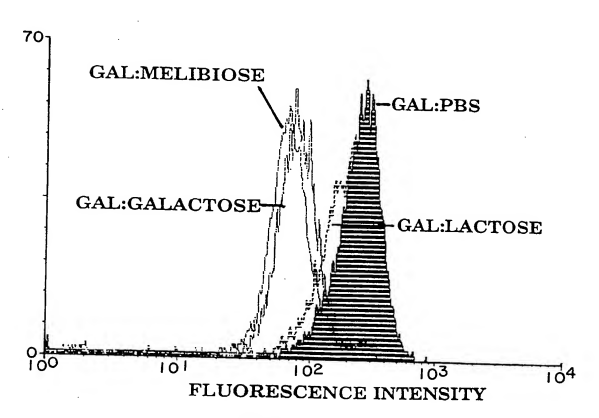


FIG. I

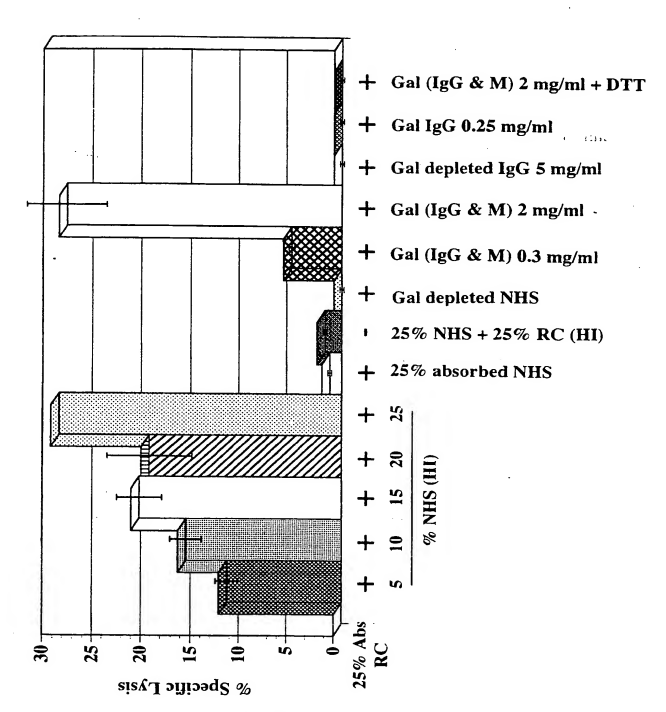
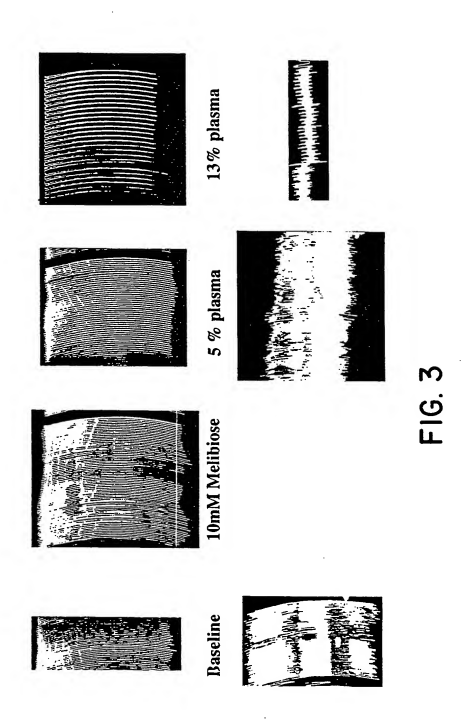


FIG. 2



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50 50 50	100 100 100	150 150 150	200 200 200	250 250 250	300 300 300	350 350 350
CAGCGCCCGC	5009505500		9999009009	c CGGATCAGGG CCTAGCCCTG	AGGAACTTGC CTTGGAATGA	Exon 2 ACGGAGTCAG TGGGAGTTGG
GCCGCTGCC	GGAGGAGCGC	CCGGCACGCC	AGCCGAGGAC	 CCAGCTTCTG TGGGTGGAGC	TGTCCCCAAG	Exon CACAAAATCA TCTCAGGCTC
GGAGCGTCGA	 ეეეემეემეე	ACACCCCGCC	CTGTTCCGGC	CCCAGCGCGC CTGGAGATTC	GCCAGCCAGC	CGAGACACTT
GGCCGAGCTG	GCCCCCTGC	GCGCAGCGGC	AGCGCGCCGA	CCGCCAGCC TCTTAGGAGG	TCCTCAAGTG	CACGGAAAGA TTCTGCTGAA
5009999900	CGGCTCCCTC	GCCGACGGGA	GGGAGGAGGC	AGCCGAGGCG	AAACCACGTG CCTTTTCTTA	CTGGCATTTG GAAGTACCGA
ਜਜਜ	51 51 51	101 101 101	151 151 151	201 201 201	251 251 251	301 301 301
PGTCD BOVGSTA MUSGLYTNS						

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400 400 400	450 450 450	500 500 500	550 550 550	009 009	650 650 650	700 700 700
CCTGCCTCCT TCTGCAGAGC CCTGCCTCCT TCTGCAGAAC CCTGCCTCCT TCGGCAGGCC	TTTTACTCTG GGGGGAGAGA TTTTACTCTA GGAGGAGAGA GTTTGCTTTG GAGGGAACAC	GAGATCTGCT TACCCCAGTC			Exon 3 Exon 4	CAATGCTGCT TGTCTCAACT CAATGCTGGT TGTCTCAACT TGATGCTGAT TGTCTCAACC
AAGGCTGCAC CTTCGCTTCC TCCCAGC	AGAGCTCACT AGAACTT-GT TTCGCC GGAGCTCAGT AGAACTT-GG TACTTTTGCC AGAAGCTCGG TTGCTTTGCT	AGCAGAGGAT GAG	TCCTGGAATT AAAGGCCTGT ACTACCTTGC	ATCACTATGC TTCAAGATCT CCATGTCAAC		TARTGRATGT CAAAGGAAGA GTGGTTCTGT CTARTGAATGT CAAAGGAAAA GTGATTCTGT TARTGAATGT CAAGGGAAAA GTAATCCTGT TSTARTGAATGT CAAGGGAAAA GTAATCCTGT TSTARTGAATGT CAAGGGAAAA
351 351 351	401 401 401	451 451 451	501 501 501	551 551 551	601 601 601	651 651 651
PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS

	707				Exon 5		
BOVGSTA	701			ATACATCAAC ATATATCCAC	AGCCCAGAAG	GTTCTTTGTT GCTCTTTGTT	750 750
MUSGLYTNS	701	. GTGGTTGTCG	TGTTTTGGGA	ATATGTCAAC	AGCCCAGACG	GCTCTTTCTT	750
PGTCD BOVGSTA MISCI VIEWS	751	CTGGATATAC	Exon 5 T CAGTCAAAAA CCCATCAAGAA		N. 4	GCTCAGAGGG AGCATTCAGA	800 800
SULITION	10/	GTGGATATAT	CACACAAAAA	TTCCAGAGGT	TGGTGAGAAC	AGATGGCAGA	800
PGTCD BOVGSTA MUSGLYTNS	801 801 801	GCTGGTGG AGGGCTGGTG AGGACTGGTG	-TTTCCGAGC GCTTCCGAGA GTTCCCAAGC	Exon TGGTTTAACA TGGTTTAACA	Exon 6 Exon 7 AACA ATGGGACTCA CAACA ATG	CAGTTACCAC GTTACCAT	850 850 850
PGTCD	851	GAAGAAGAAG	ACGCTATAGG	CAACGAAAAG		AAGAAGACAA	006
BOVGSTA	851	GAAGAAGATG	GAGACATAAA	CGAAGAAAAG	•	ACGAAGACGA	006
MUSGLYTNS	851	GAAGACAACG	TAGAAGGACG	GAGAGAAAAG	GGTAGAAATG	GAGATCGCAT	006
PGTCD	901	CAGAGGAGAG	CTTCCGCTAG	Դ ՀՀՋ ՐԳՀՀԳԳ	Exon 7	Exon 8	0.20
BOVGSTA	901	AAGCAAG	CTTAAGCTAT	CGGACTGGTT		AAACGCCCG	950
MUSGLYTNS	901	TGAAGAG	CCTCAGCTAT	GGGACTGGTT	•	AACCGCCCGG	950
PGTCD		AGGTCGTGAC	CATAACCAGA	TGGAAGGCTC	CAGTGGTATG G	GGAAGGCACT	1000
BOVGSTA	951	AGGTTGTGAC	CATGACGAAG	TGGAAGGCTC	CAGTGGTGTG G	GGAAGGCACT	1000
MUSGLYTMS		ATGTTTTGAC	AGTGACCCCG	TGGAAGGCGC	CGATTGTGTG G	GGAAGGCACT	1000
	1001	TACAACAGAG	CCGTCTTAGA	TAATTATAT	GCCAAACAGA A	AAATTACCGT	1050
	1001	TACAACAGAG	CCGTCTTAGA	CAATTATTAT	GCCAAGCAGA A	AAATTACCGT	1050
MUSGLYTMS	1001	TATGACACAG	CTCTGCTGGA	AAAGTACTAC	GCCACACAGA A	AACTCACTGT	1050
							1

PGTCD BOVGSTA	1051 1051	GGGCTTGACG	GTTTTTGCTG	Exon 8 Exon 9 TCGGAAGATA CAT	xon 9 CATTGAGCAT CATTGAGCAT	TACTTGGAGG	1100
MUSGLYTNS	1051	GGGGCTGACA	GTGTTTGCTG			_	1100
BOVGSTA MUSGLYTNS	1101	AGTTCTTAAC ACTTTCTGGA	AICIGCAAI GTCTGCTAAT GTCTGCTGAC	ACATACTTCA AAGCACTTCA ATGTACTTCA	TGGTTGGCCA TGGTGGGCCA TGGTTGGCCA	CAAAGTCATC CCCAGTCATC TCGGGTCATA	1150 1150 1150
PGTCD	1151	TTTTACATCA	TGGTGGATGA	TATCTCCAGG	ATGCCTTTGA	TAGAGCTGGG	1200
BOVGSTA	1151	TTTTATATCA	TGGTAGATGA	TGTCTCCAGG	ATGCCTTTGA	TAGAGTTGGG	1200
MUSGLYTNS	1151	TTTTACGTCA	TGATAGATGA	CACCTCCCGG	ATGCCTGTCG	TGCACCTGAA	1200
PGTCD	1201	TCCTCTGCGT	TCCTTTAAAG	TGTTTGAGAT	CAAGTCCGAG	AAGAGGTGGC	1250
BOVGSTA	1201	TCCTCTGCGC	TCCTTCAAAG	TGTTTAAGAT	CAAGCCTGAG	AAGAGGTGGC	1250
MUSGLYTNS	1201	CCCTCTACAT	TCCTTACAAG	TCTTTGAGAT	CAGGTCTGAG	AAGAGGTGGC	1250
PGTCD	1251	AAGACATCAG	CATGATGCGC	ATGAAGACCA	TCGGGGAGCA	CATCCTGGCC	1300
BOVGSTA	1251	AGGACATCAG	CATGATGCGC	ATGAAGACTA	TCGGGGAGCA	CATTGTGGCC	1300
MUSGLYTNS	1251	AGGATATCAG	CATGATGCGC	ATGAAGACCA	TTGGGGAGCA	CATCCTGGCC	1300
PGTCD	1301	CACATCCAGC	ACGAGGTGGA	CTTCCTCTTC	TGCATGGACG	TGGATCAGGT	1350
BOVGSTA	1301	CACATCCAGC	ATGAGGTTGA	CTTCCTTTTC	TGCATGGATG	TGGACCAGGT	1350
MUSGLYTNS	1301	CACATCCAGC	ACGAGGTCGA	CTTCCTCTTC	TGCATGGACG	TGGATCAAGT	1350
PGTCD BOVGSTA MUSGLYTNS	1351 1351 1351	CTTCCAAAAC Z	AACTTTGGGG AAGTTTGGGG	TGGAGACCCT TGGAGACCCT TGGAAACTCT	GGGCCAGTCG GGGCGAGTCG GGGCCAGCTG	GTGGCTCAGC GTGGCCCAGC GTAGCACAGC	1400 1400 1400

CTACGAGAGG 1450	ATTTTTATTA 1500	AACATCACTC 1550	CATAGAAGCC 1600	TCAACAAACC 1650	ATAGGCATGT 1700	AGAGTATAAT 1750
CTACGAGAGG 1450	ATTTTTATTA 1500	AACATCACCC 1550	CATAGAAGCC 1600	TCAACAAACC 1650	ATAGGCCTAC 1700	AGAGTATAAT 1750
CTATGAGAGG 1450	ATTTTTACTA 1500	AACCTCACCA 1550	CATAGAAGCC 1600	TCAACAAACC 1650	ATAGGCCTGC 1700	AGAGTATAAT 1750
ACGAGTTCAC	GGCCAGGGGG	TCAGGTTCTA	AGGAAAATGA	TATTTAATTC TO	GGATTATCAT AT	GGCAGAAAAA AG
ATGACTTCAC	GGCGAAGGGG	TCAGGTCCTT	AGAAAAATGA	TATTTCCTTC TO	GGATTATCAC AT	GGCAGACAAA AG
AGAAGTTCAC	GGAGAGGGGG	TCACATTCTC	AGAAACATGA	TACTTCCTTT TO	GGACTATCAG AT	GGCAGACAAA AG
GCACATCCTG GCAGATCCCA GCCAGTCCCG	CATTCCGTTT CATTCCCTTC	GAACACCCAC GAACACCCAC GAACGCCTAC	CTCCAGGACA CTCAAGGACA CTCCAGGACA	TCTAAACAAG TCTAAACAAG CCTCAACAAA	AATACTGCTG AATACTGCTG AGTATTGCTG	AAGATAGCTT AAGATGTCTT AAGGTAGCTT
GTGGTACAAG	CCGCAGCCTA	ATTTTTGGGG	CAAGGGAATC	ATGAAAGCCA	TTATCCCCAG	TAGGATTGTC AL
GTGGTACAAG	CTGCAGCATA	ATTTTTGGGG	CAAAGGAATC	ATGAAAGCCA	TTATCCCCGG	TAAGCTTGTC AL
GTGGTACAAG	CGGCCGCGTA	ATTTTTGGAG	TAAGGGGATC	ATGAGAGCCA	CTATCTCCAG	TAAAAGTGTC AA
TACAGGCCTG	CGGAAGGAGT	CCACGCAGCC	AGGAGTGCTT	GAGTGGCATG	CACTAAAATC	CTGTGGATAT
TACAAGCCTG	CGGAAGGAGT	CCATGCAGCC	AGGAATGCTT	CAATGGCATG	TACTAAAATC	CTGCGGATAT
TCCAGGCCTG	CGGGAACTGT	CCACGCGGCC	GGGAGTGCTT	CAGTGGCATG	CACTAAAATC	CTTCAGATAT
1401	1451	1501	1551	1601	1651	1701
1401	1451	1501	1551	1601	1651	1701
1401	1451	1501	1551	1601	1651	1701
PGTCD	PGTCD	PGTCD	PGTCD	PGTCD	PGTCD	PGTCD
BOVGSTA	BOVGSTA	BOVGSTA	BOVGSTA	BOVGSTA	BOVGSTA	BOVGSTA
MUSGLYTNS	MUSGLYTNS	MUSGLYTNS	MUSGLYTNS	MUSGLYTNS	MUSGLYTNS	MUSGLYTNS

1751 TTGGTTAGAA ATAACATQTG ACTTTAAATT GTGCCAGCAG TTTTCTGAAT 1800 a 1751 GTGGTTAGAA ATAATGTQTG ACTTT GTGCCAGTAC ATTTCTGAAT 1800 INS 1751 TTGGTTAGAA ATAATGTQTG ACTTCAAATT GTGATGGAAAC 1800 Stop	1801 TTGAAAGAGT ATTACTCTGG CTACTTCCTC AGAGAAGTAGCACTTAA 1850 A 1801 TTGAGAGAGT ATTATTCTGG CTACTTCCTC AGAAAAGTAACACTTAA 1850 INS 1801 TTGACACT ATTACTCTGG CTAATTCCTC AAACAAGTAG CAACACTTGA 1850	1851 TTTTAACTTT TAAAAAATA CTAACAAAATACCAA CACAGTAA-G 1900 1 1851 TTTTAACTTA AAAAAAATA CTAACAAAAGACCAA CACAGCAA-A 1900 NS 1851 TTTCAACTTT TAAAAGAA-A CAATCAAAAC CAAAACCCAC TACCATGGCA 1900	1901 TACATATTAT TCTTCCTTGC AACTTTGAGC CTTGTCAAAT GGGAGAATGA 1950 1901 TACATATTAT TTCTCCTTGT AACTTTGAGC CTTGTAATAC GGGAGAATGA 1950 NS 1901 AACAGATGAT TTCTCCT-GA CACCTTGAGC CT-GTAATAT GTGAGAAAGA 1950	1951 CTCTGTGGTAATCAGA TGTAAATTCC CAGTGATTTC 2000 1951 ACCTGTGGTAATCAGA TGTAAATTCC CAGTGATTTC TTACCTATTT 2000 NS 1951 GTCTATGGCA AGTAATCAGG TATAAATTCT CAATGATTTC TTATATATTC 2000	2001	2051
1751 1751 1751	1801 1801 1801	1851 1851 1851	1901 1901 1901	1951 1951 1951	_	
. PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS	PGTCD 2 BOVGSTA 2 MUSGLYTNS 2

PGTCD BOVGSTA MUSGLYTNS	2101 2101 2101	GGCCAGGTGC	TGAGAGAAGC	ATTAGGGAAC	AGTGTGGGTT	GTGTCAGAGT	2150 2150 2150
PGTCD BOVGSTA MUSGLYTNS	2151 2151 2151	TGGACGGCTC	CATCCCTTTG	GCTTCATTAT	CTTCCTCCTC	ATGGAGATTC	2200 2200 2200
PGTCD BOVGSTA MUSGLYTNS	2201 2201 2201	TAAAGCAACC	CAGAGAGGCT	TTGCAGCCAG	AGACCTTTAA	TAAGGATGCC	2250 2250 2250
PGTCD BOVGSTA MUSGLYTNS	2251 2251 2251	AATGTGACCA	TCAGTCTGTA	AAAGCTGATG	GCTCCAGGAG	CGCTGGCAGT	2300 2300 2300
PGTCD BOVGSTA MUSGLYTNS	2301 2301 2301	CCAGGCCCCA	CTAGGCTATT	GITTCTGTCC	TGGGCATAAA	GGAGGCAGAG	2350 2350 2350
PGTCD BOVGSTA MUSGLYTNS	2351 2351 2351	AGTGCCAATA	GGTACTTTGG	TGGCACATGT	TCAGAGTCCA	GGAAAAATCA	2400 2400 2400
PGTCD BOVGSTA MUSGLYTNS	2401 2401 2401	AGGGTGACCA	CTTAGAGGGA	сатаддастт 4 0	GGGGTTGGTG	ATTGAACTGA	2450 2450 2450 2450

2500 2500 2500	2550 2550 2550	2600 2600 2600	2650 2650 2650	2700 2700 2700	2750 2750 2750	2800 2800 2800
AGGAATTGAA	TGCTGTTAGC	GGGAAGGAGA	AAGACAATCT	GAGTTCCTCA	GACTGAAGAC	ACTATGCAAG
GACTAACAGC	ATTGTATTCA	GACTGTATCA	ATTATATCAG	CTGAACCTCC	TGGAGCAGGT	CTGCTCTAGT
TCTTCAGGAT	TTTTGCCCAA	GAGAGGGTGT	CCAGCACCCT	ACAACCTGCT	GGGCCTGTA	CTCATTTCCC
CAGACAGCTT	GTTCATTTG	GAGCCCTGTG	GGACTGAGGA	GGTCCTACCT	TTCCAGTGTG	GTCACATGAC
GTTACAAACA	TGGAAAGTGT	TTTGTGTGTT	GTACCTCAGC	CTCATCATCA	GCCCATCGTG	AAAGCCCCCT
2451 2451 2451	2501 2501 2501	2551 2551 2551	2601 2601 2601	2651 2651 2651	2701 2701 2701	2751 2751 2751
PGTCD BOVGSTA MUSGLYTNS						

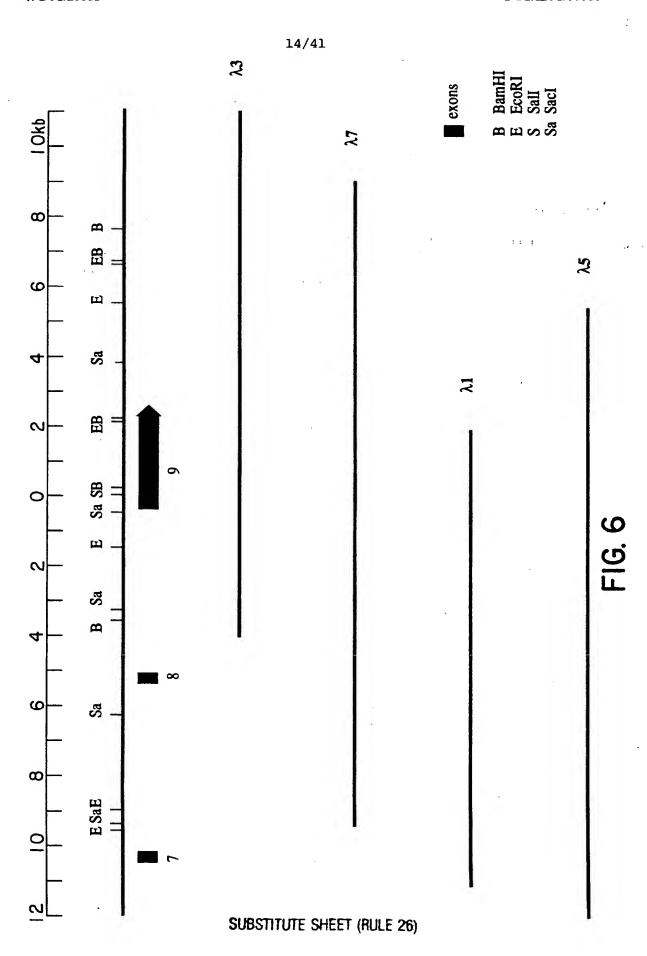
2850 2850 2850	2900 2900 2900	2950 2950 2950	3000 3000 3000	3050 3050 3050	3100 3100 3100
TTT	GTT			GTAACAGTAG	GTC
ACCGACTTT	GGTTCTGTT	CTCAACCTA	CAGGTGTCC	GTAACAGTA	GGTTCGTGT
ACC	GGT	···		GTA	GGTTCGTGTC
CAACATAGGA ACCGACTITA	AGCTGCTGAA GGTTCTGTTC		G	G.	
AACATAGGA	GCT	GTG		rgac	
CAACATAGGA	AGCTGCTGAA	ATAGGTGGTT	GACCCTTTCA	ACTCTGACTG	ATAATTCTGG
		Ł			A
ATGTACTGGA	ACTACAACGG	GAGCCCCTGT	AGTGTTAAAT	AGATATTTCC	 AGCAAGGGAA
 GTA	TAC		TGT	ATAC	CAAC
		: : &			A
AGCCAGCCAG	AGCCGCAGTC	GAGCCTGCAG GAGCCCCTGT	CCTTTGGGA	TAAAAAACAT	 AAT
CAG		CCTC	CCTTTGGGA	AAA	GTTATGAAAT
AGC	AGC	GAG	CCCTTTGGGA	TAAAAAACAT	GTTATGAAAT AGCAAGGGAA ATAATTCTGG
	ggg	rg.	3.A.C		
ACA(AAT	CHC	3000	TAAGACGG	TTA
TGTGACAGCC	TGGCAATGGG	CCCGCTCTGA	GGGTCGCGAC	CTAAGACGGT	CAGAATTACA
2801 2801 2801	2851 2851 2851 2851	2901 2901 2901	295 295 295	3001 3001 3001	3051 3051 3051
A TNS	A I'NS	a Ins	A INS		
PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS	PGTCD BOVGSTA MUSGLYTNS
PGTCD BOVGS MUSGL	PGTCD BOVGS MUSGL	PGTCD BOVGS MUSGL	PGTCD BOVGS MUSGL	PGTCD BOVGS MUSGL	PGTCD BOVGS' MUSGL

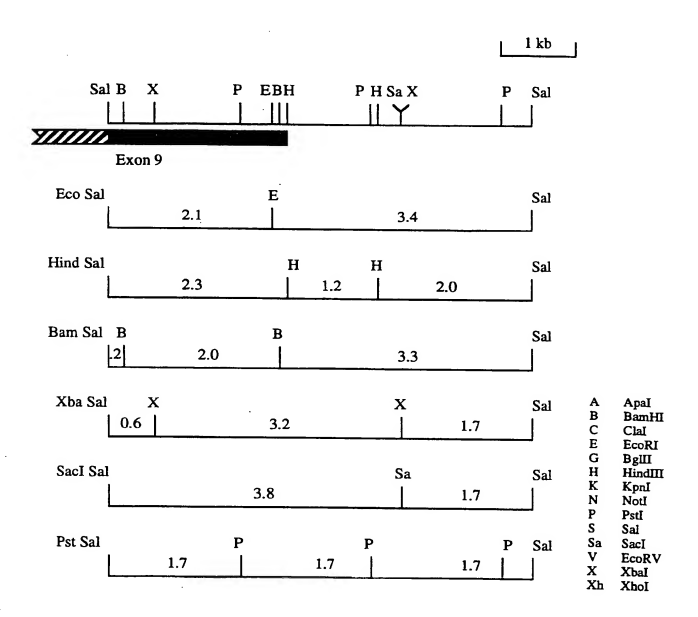
F16. 4

					Ex4 VEx	Ex5 VE	3 ×6	
PGT[Frame			MNVKGRVVLS	MLLVSTVMVV	FWEYINSPEG	SLEWIYOSKN	PEVC-CCAOD	5(
BGT[Frame	1)	1	MNVKGKVILS	MLVVSTVIVV	FWEYTHSPEG	ST.FWINDSDN	DEVICECTOR	E (
MGT[Frame	1]	1	MNVKGKVILL	MLIVSTVVVV	FWEYVNSPDG	SFLWIYHTKI	PEVGENRWQK	5(
				Ex7 .				
PGT[Frame	1]	51	GWWFPSWFNN	GTHSYHERED	ATCHEREODE	EDNRGELPLV	Ex7 Ex8	101
BGT[Frame	1]	51	GWWLPRWFNN	GYHEEDG	DINEEKEODN	ED-ESKLKLS	DWFNPEKRPE	100
MGT[Frame	1]	51	DWWFPSWFKN	GTHSYOEDNV	EGRREK-GRN	GDRIEEPQLW	DMENDAMDDD	100
				_				100
DCT [Exame	7 14	^-				Ex	8 E x9	
PCT[Frame	117	.01	VVTITRWKAP	VVWEGTYNRA	VLDNYNAKQK	ITVGLTVFAV	GRYIEHYLEE	150
DGILLIUME	T 1 T	UL	VVIMIKWKAP	VVWEGTYNRA	VI.DNVVX KOV	TONIOT MITTARY	COULTBURE DE	
rigitiame	T 1 T	ΟŢ	VLTVTPWKAP	IVWEGTYDTA	LLEKYYATQK	LTVGLTVFAV	GKYIEHYLED	150
PGT[Frame	1]1	51	FLISANTYFM	VGHKVIFYIM	VDDISRMPLI	ELGPLRSFKV	FFTKGEKDWO	200
DGT [LT GIME	TIT	2.T	FLTSANKHFM	VGHPVIFYTM	VDDVSPMPI.T	FI CDI DCDVV	DUTUDDUDO	
MGT[Frame	1]1	51	FLESADMYFM	VGHRVIFYVM	IDDTSRMPVV	HLNPLHSLQV	FEIRSEKRWO	200
								200
PGT[Frame	1]2	01	DISMMRMKTI	GEHILAHIOH	EVDELECMDV	DQVFQNNFGV	Emt cogus or	
DOLLILAMO	+ 1 4	OI.	DISUMBUKLI	GEHIVAHTOH	EVIDELECMOV	DOMEODADOM	DMY 070000	0 - 0
MGT[Frame	1]2	01	DISMMRMKTI	GEHILAHIOH	EVDFLFCMDV	DQVFQDNFGV	ETLGESVAQL	250
				_	, , , , , , , , , , , , , , , , , , ,	DQ II QDMFGV	FILEGUANGE	250
PGT[Frame	112	51	OAWWYKAHPD	FFTVFDDVFC	AAVIDEGGG	FYYHAAIFGG		
BGT[Frame	112	51	OAWWYKADPN	DETVEDENCES	AATIPEGGGD	FYYHAA1FGG FYYHAA1FGG	TPTQVLNITQ	300
MGT[Frame	1]2	51	OAWWYKASPE	KFTVFDDFIC	AATIPFGEGD	FYYHAAIFGG FYYHAAIFGG	TPTQVLNITQ	300
		_	2	MI I I EKKEUS	AATIPFGEGD	FYYHAAIFGG	TPTHILNLTR	300
PCT (Frame	113	0.3	Dobugar care					
RGT[Frame	112	01	ECFKGILQDK	ENDIEAEWHD	ESHLNKYLIL	NKPTKILSPE	YCWDYHIGMS	350
oo - [- Lame	T 1 3	O T	ECLUGITUDE	KNDIFAOWHD	FCHI MVVETT	MIV DOUGHT COO		_
Joiliame	1 13	υŢ	ECFKGILQDK	KHDIEAQWHD	ESHLNKYFLF	NKPTKILSPE NKPTKILSPE	YCWDYQIGLP :	350
D.O					•			
rGT Frame	1]3!	51	VDIKIVKIAW	QKKEYNLVRN	NI*	•••••		.00
	- ,		UD TIVID A VIJIDA	UIRP.YNVVRN	MV ×			00
MGT Frame	1]3	51	SDIKSVKVAW	QTKEYNLVRN	NV*	• • • • • • • • • • • • • • • • • • • •	4	00

FIG. 5

WO 95/20661 PCT/IB95/00088



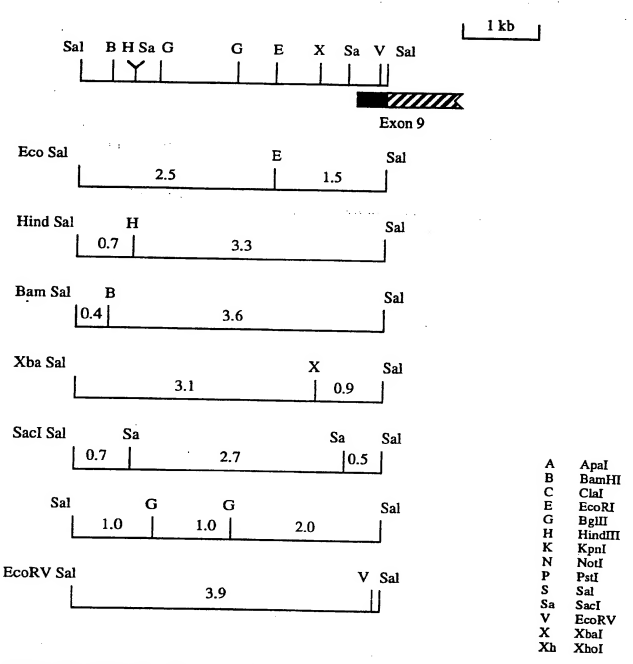


No sites for: BglII, Nde, PvuI, Xho, Kpn, SacII, EcoRV, Sma, Cla, Apa, Not

pBS+KS: SacI SacII Not Xba Spe Bam Sma Pst Eco RV Hind Cla Sal Xho Apa Kpn

1

FIG. 7



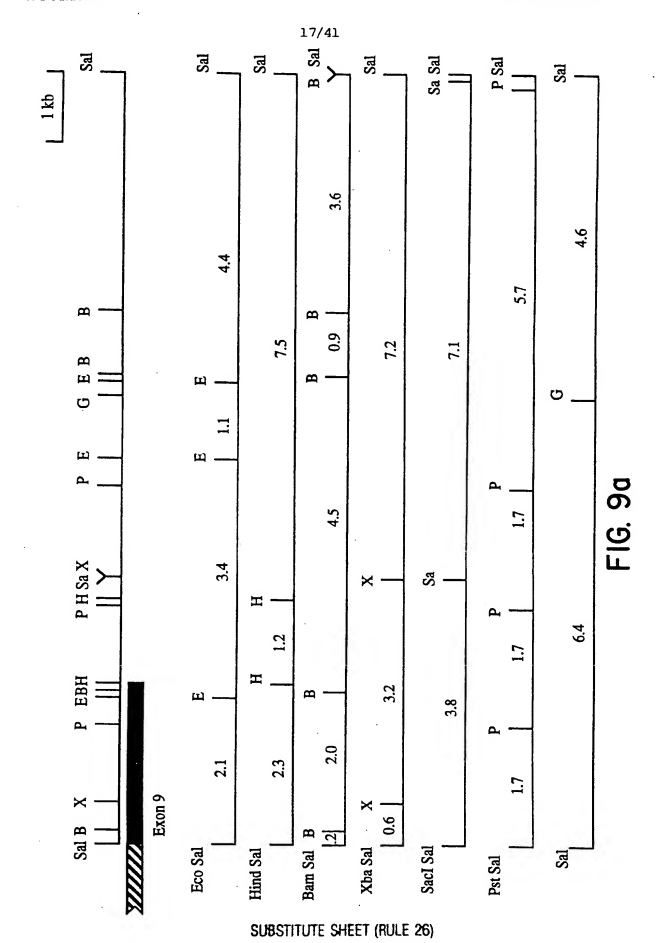
No sites for: Nde, PvuI, Xho, Kpn, SacII, Sma, Cla, Apa, Not

Unmapped sites for: Pst, PvuII

pUBS:

.... SacI SacII Not Xba Spe Bam Sma Pst Eco RV Hind Cla Sal Xho Apa Kpn

FIG. 8



Apal
BamHI
Clal
Clal
EcoRI
BglII
HindIII
KpnI
Notl
Pstl
Sal
Sacl
EcoRV
Xbal XX < SS O P Z X H G E C B A

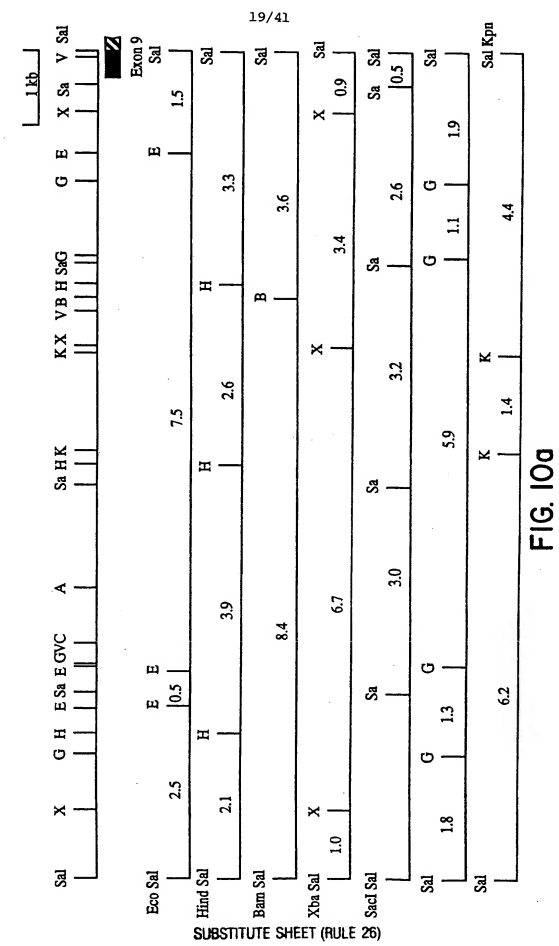
FIG. 9b

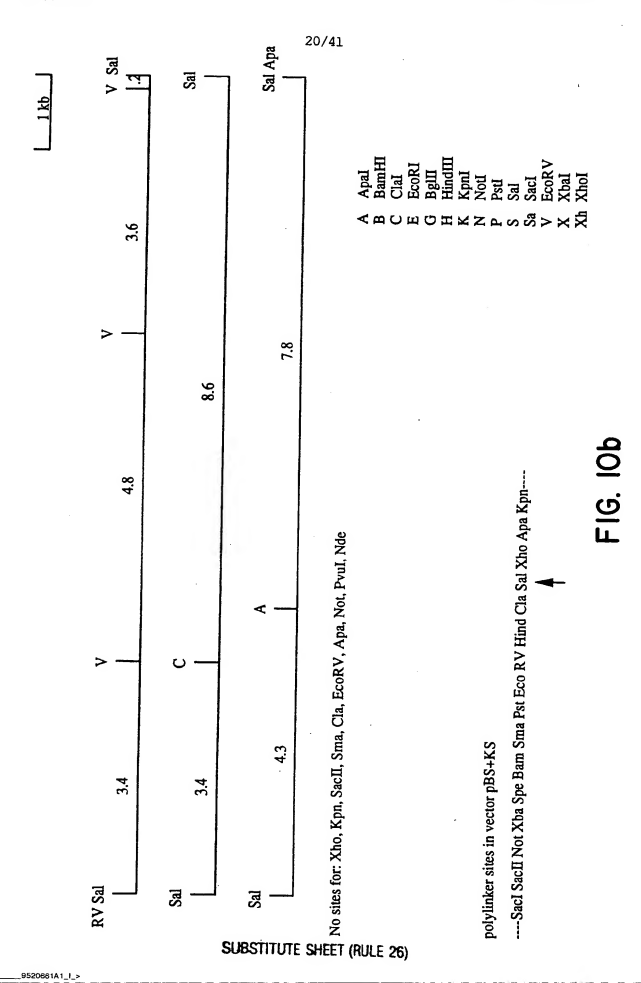
polylinker sites in vector pUBS (pUC19 with polylinker from pBluescript M13+):

No sites for: Xho, Kpn, SacII, Sma, Cla, EcoRV, Apa, Not, PvuI, Nde

.... SacI SacII Not Xba Spe Bam Sma Pst Eco RV Hind Cla Sal Xho Apa Kpn

PCT/IB95/00088





3NSDOCID: <WO_____9520661A1_I_>

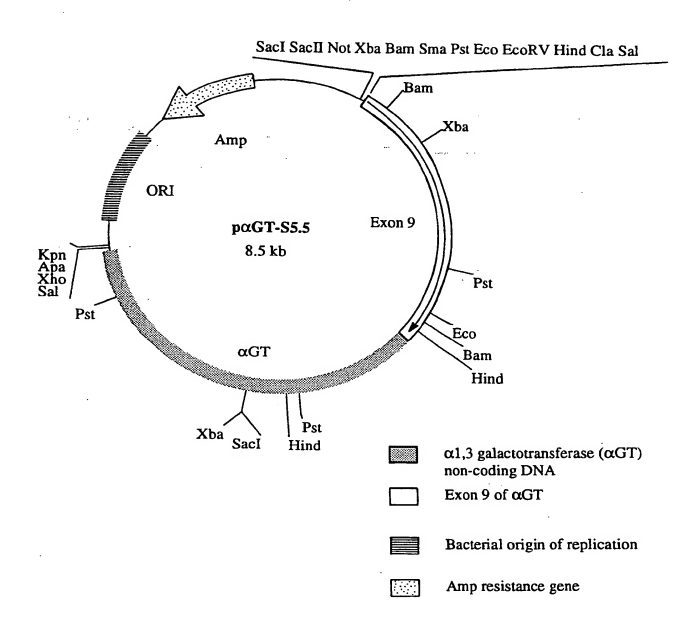


FIG. II

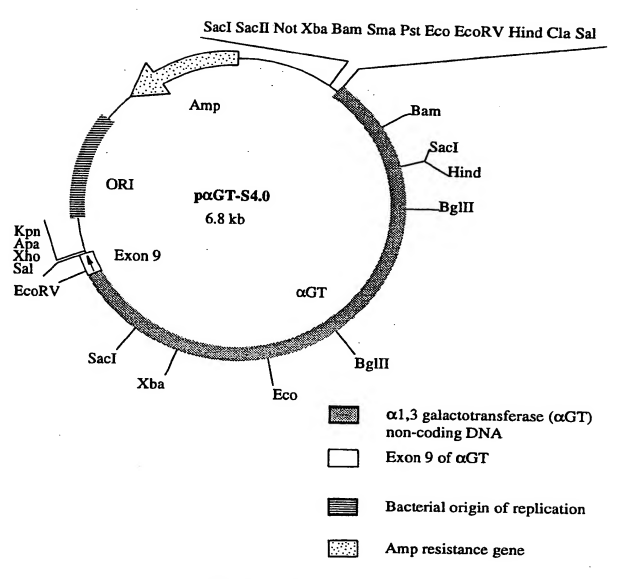
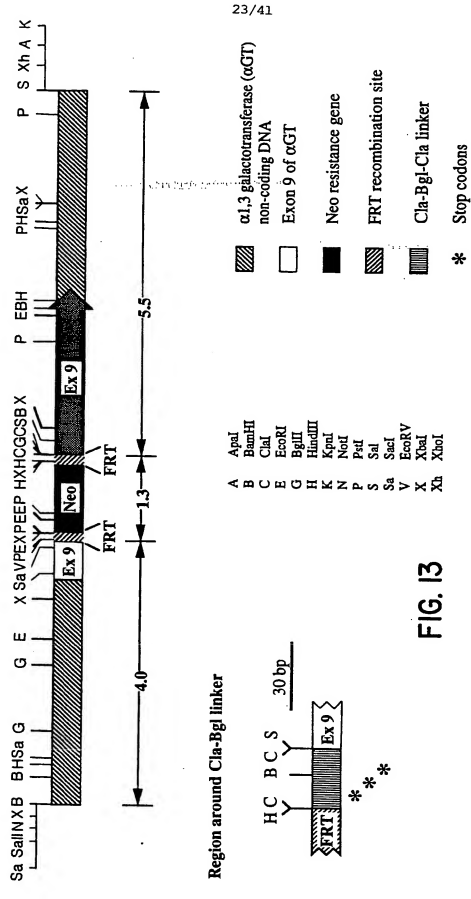
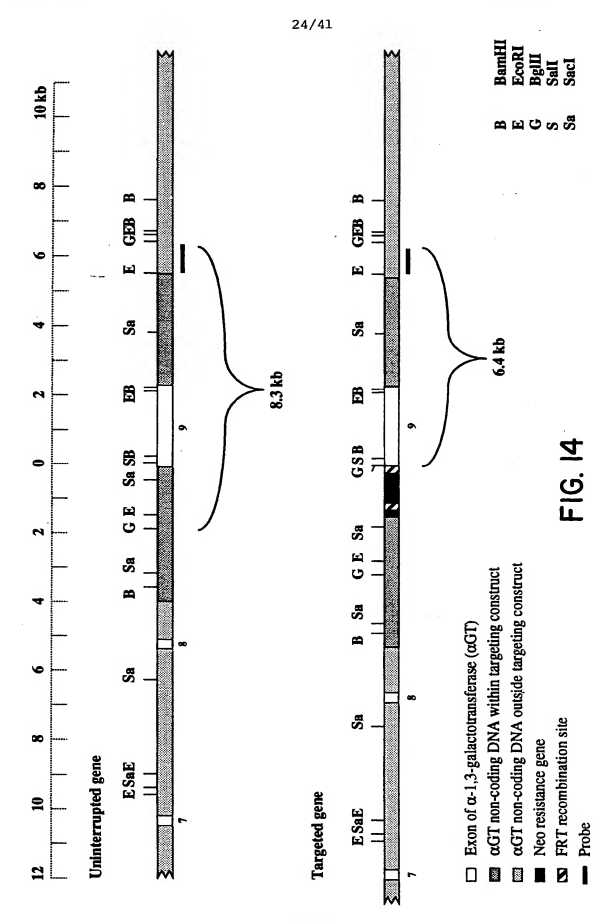


FIG. 12

3NSDOCID: <WO_____9520661A1_I_>

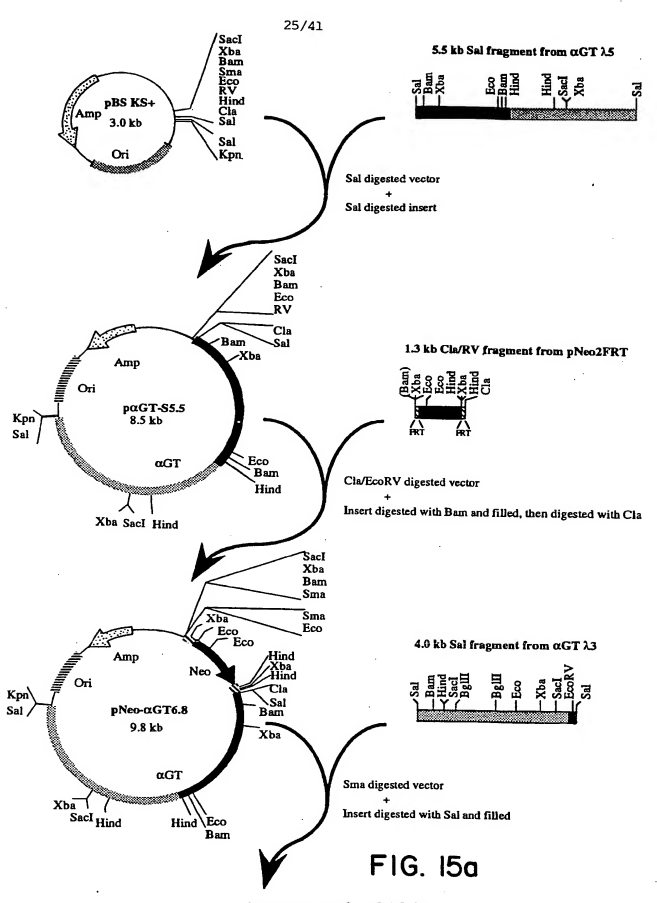


SUBSTITUTE SHEET (RULE 26)



SUBSTITUTE SHEET (RULE 26)

PCT/IB95/00088



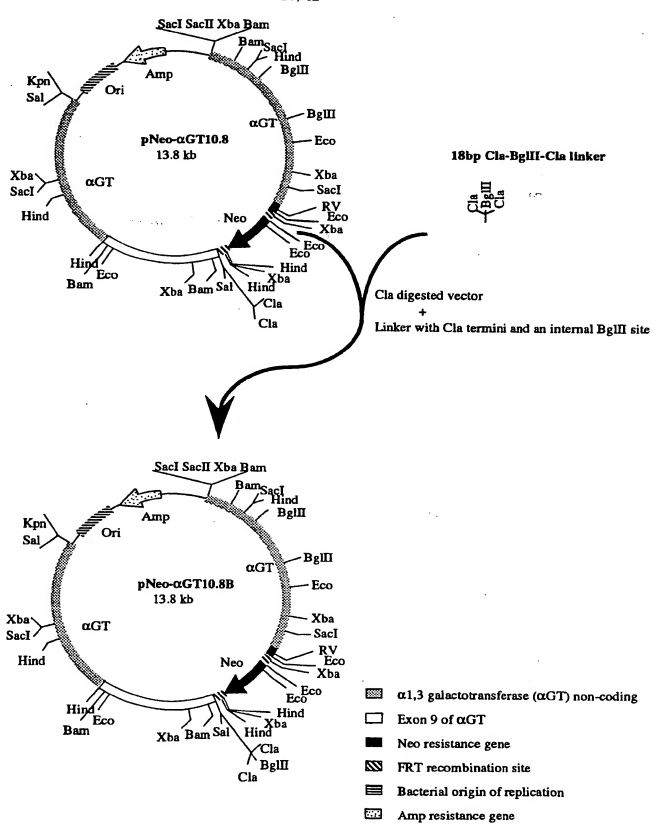


FIG. 15b

60 GTTCCTATTC	50 (T) TTTCCC > >	40 CTTGA ACTTC	30 GATCCCCCAG	20 GGAATTCGAT	10 GAGGGCTGCA
· ·					
120	110	100	-: 90	80	70
120 AGTGTGGTTT					
180	170	160	150	140	130
180 CAATGTCGAG 240	TGTTTCCACC	AGGCCTGGAA	CTCTCCACCC	AGCAAAAAGC	TGCAAGAGGA
240	230	220	210	200	190
ATGTTTCCAC	CAGGCCTGGA	CCTCTCCACC	AAGCAAAAAG	TTGCAAGAGG	CAGTGTGGTT
					250
300	290	280	270	260	
ACGCAGATGC	GAATTCGAAC	TGTCATTGGC	CCCAGCGTCT	GCAAACCCCG	CCAATGTCGA
360	350	340	330	320	310
GTGGCCTCGA	330	CCOMAMANA	ACCTICCACTUTE	GCGCGGTCCC	AGTCGGGGCG
420	410	400	390	380	370
ATTGCACGCA	210	TCCCCC A TOTO	СУУФУЛСССУ	ACCCTGCAGC	ACACCGAGCG
480	470	460	450	440	430
ACAGACAATC	ACTGGGCACA	TTCGGCTATG	GGAGAGGCTA	CCGCTTGGGT	GGTTCTCCGG
540	530	520	510	500	490
TCTTTTTGTC	GGCGCCCGGT	TCAGCGCAGG	GTTCCGGCTG	ATGCCGCCGT	GGCTGCTCTG
600	590	580	570	560	550
	ACCCACCCCC	CTCCAAGACG	CCTGAATGAA	TGTCCGGTGC	AAGACCGACC
660	650	640	630	620	610
AGCGGGAAGG	TTGTCACTGA	GTGCTCGACG	TTGCGCAGCT	CGGGCGTTCC	CTGGCCACGA
				680	670
720	/10	700	0 E O	TATION COCK	
		CAGGATCTCC			
780	770	760	750	740	730
TC MCCCCCC	TCC ATTACCOT	ATGCGGCGGC	GGCTGATGCA	TATCCATCAT	GCCGAGAAAG
840	830	820	810	800	790
TCGGATGGAA	GAGCACGTAC	CGCATCGAGC	AGCGAAACAT	TCGACCACCA	ACCTGCCCAT

FIG. 16a

850	860	870	880	890	900
GCCGGTCTTG	TCGATCAGGA	TGATCTGGAC	GAAGAGCATC	AGGGGCTCGC	GCCAGCCGAA
910	920	930	940	950	960
CTGTTCGCCA	GGCTCAAGGC	GCGGATGCCC	GACGGCGAGG	ATCTCGTCGT	GACCCATGGC
970	980	990	1000	1010	1020
GATGCCTGCT	TGCCGAATAT	CATGGTGGAA	AATGGCCGCT	TTTCTGGATT	CATCGACTGT
1030	1040	1050	1060	1070	1080
GGCCGGCTGG	GTGTGGCGGA	CCGCTATCAG	GACATAGCGT	TGGCTACCCG	TGATATTGCT
1090	1100	1110	1120	1130	1140
GAAGAGCTTG	GCGGCGAATG	GGCTGACCGC	TTCCTCGTGC	TTTACGGTAT	CGCCGCTCCC
on rooms	1160 GCATCGCCTT	CTATCGCCTT	CTTGACGAGT	TCTTCTGAGG	GGATCGGCAA
	1220 GAATAAAACG	CACGGGIGII	GGGCGTTTTGT	TCGGATCATC	AAGCTTGAAG
	1280 GAAGTTCCTA	TTCTCTAGAA	AGTATAGGAA	CTTCAAGCTT	ATCGATGAGT
1330 AGATCTTGAT	1340 CGATACCGTC	1350	1360	1370	. 1380

Linker sequences: 0-28

FRT: 29-104

Polyoma virus enhancer repeats: 105-249

Herpes Simplex Virus Tyrosine Kinase promoter: 250-385 Neomycin phosphotransferase coding region: 385-1188

Herpes Simplex Virus Tyrosine Kinase PolyA signal: 1189-1249

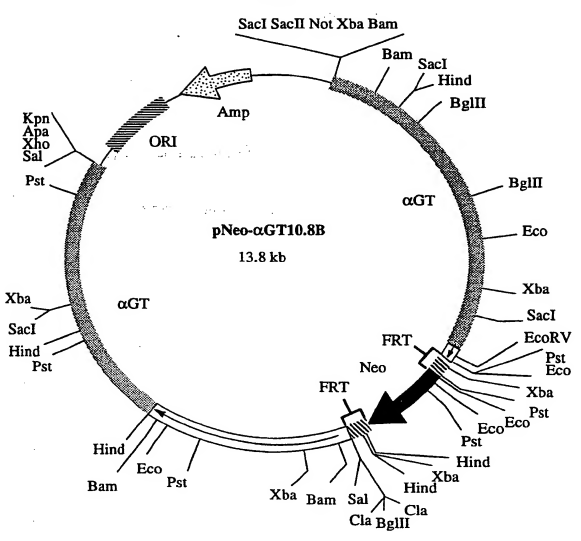
FRT: 1250-1310

Linker sequences: 1311-1340

FIG. 16b

PCT/IB95/00088





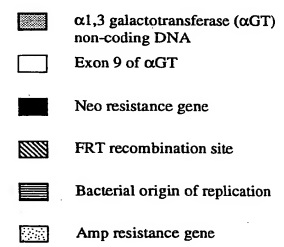
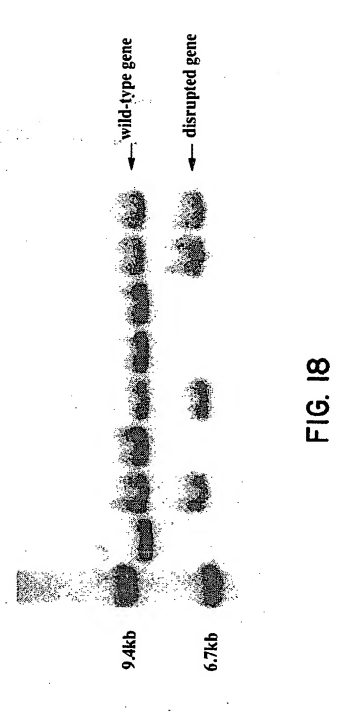
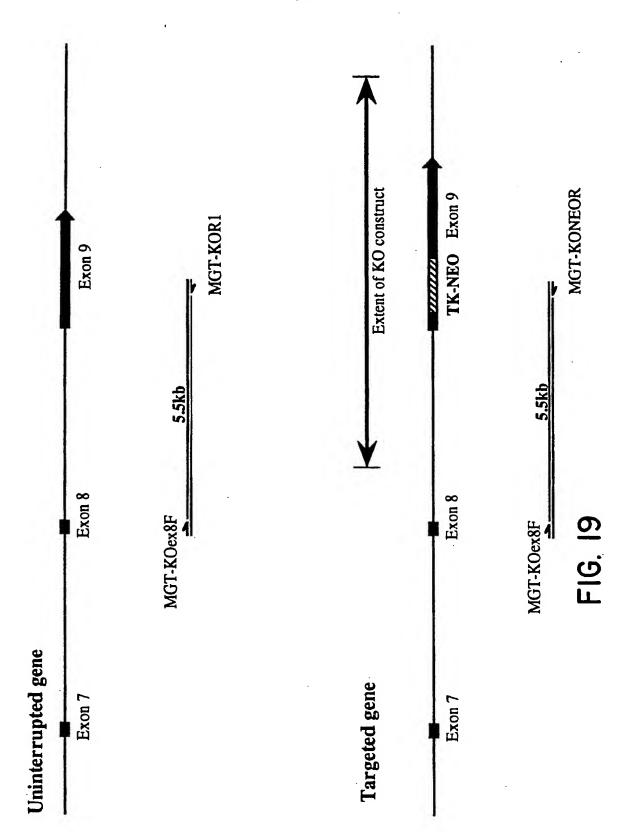


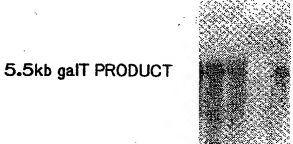
FIG. 17



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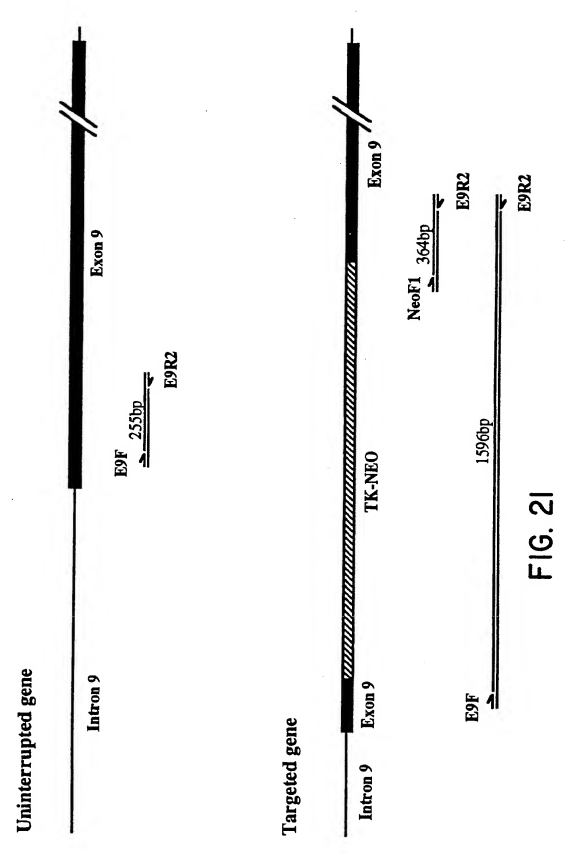


5.5kb KO PRODUCT

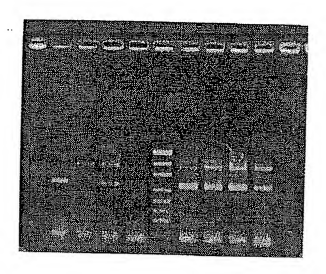
- 1. CBAC TEMPLATE; WILD TYPE PRIMERS
- 2. 7C2 TEMPLATE; WILD TYPE PRIMERS
- 3. CBAC TEMPLATE; KO PRIMERS
- 4.7C2 TEMPLATE; KO PRIMERS

FIG. 20





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364bp 255bp

FIG. 22

9R2 983bp cDNA spike product Primer binding sites within mouse ferrochelatase cDNA Primer binding sites within mouse a.1,3-GalT cDNA FIG. 23 Exons

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MCKHLKHL MKHL

i) Ferrochelatase, FC-F/R

M, Marker SPP-I C, MQW control K, KIDNEY H, HEART L, LIVER

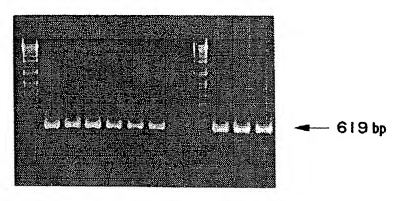
FIG. 24a

→ 709 bp

<u>+/- +/+</u> мкнг

ii) α-1,3-GT cDNA spike + 7F/9R2 primers

FIG. 24b

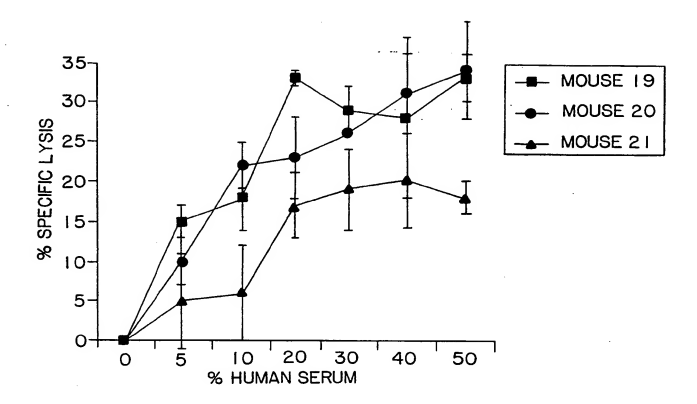


M C K H L K H L M K H L

iii) α-1,3-GT 7F/9R2 primers

FIG. 24c

→ 619 bp



MOUSE 19: WILD TYPE; MOUSE 20: HETEROZYGOTC Gal KO; MOUSE 21: HOMOZYGOUS Gal KO

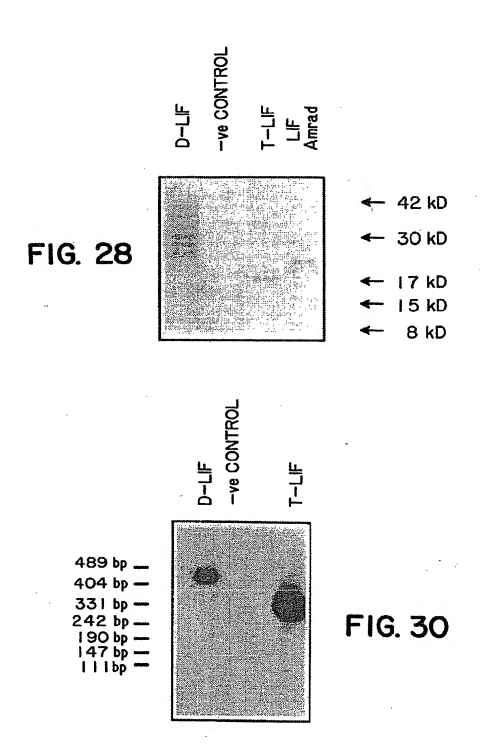
FIG. 25

60	GGAGTCCAGCCCATAATGAAGGTCTTGGCCGCAGGGATTGTGCCCCTTACTGCTGCTGGTT	1
120	CTGCACTGGAAACACGGGGCAGGGAGCCCTCTTCCCATCACCCCTGTAAATGCCACCTGT	61
10	MetAsnGlnIleLysAsnGlnLeuAlaGln	1
180	GCCATACGCCACCCATGCCACGCCAACCTCATGAACCAGATCAAGAATCAACTGGCACAG	121
30	Leu Asn Gly Ser Ala Asn Ala Leu Phe Ile Ser Tyr Tyr Thr Ala Gln Gly Glu Pro Phe	10
240	CTCAATGGCAGCGCCAATGCTCTCTTCATTTCCTATTACACAGCTCAAGGAGAGCCGTTT	181
50	ProAsnAsnValGluLysLeuCysAlaProAsnMetThrAspPheProSerPheHisGly	30
300	CCCAACAACGTGGAAAAGCTATGTGCGCCTAACATGACAGACTTCCCATCTTTCCATGGC	241
70	AsnGlyThrGluLysThrLysLeuValGluLeuTyrArgMetValAlaTyrLeuSerAla	50
360	AACGGGACAGAGAGACCAAGTTGGTGGAGCTGTATCGGATGGTCGCATACCTGAGCGCC	301
90	SerLeuThrAsnIleThrArgAspGlnLysValLeuAsnProThrAlaValSerLeuGln	70
420	TCCCTGACCAATATCACCCGGGACCAGAAGGTCCTGAACCCCACTGCCGTGAGCCTCCAG	361
110	ValLysLeuAsnAlaThrIleAspValMetArgGlyLeuLeuSerAsnValLeuCysArg	90
480	GTCAAGCTCAATGCTACTATAGACGTCATGAGGGGCCTCCTCAGCAATGTGCTTTGCCGT	421
130	LeuCysAsnLysTyrArgValGlyHisValAspValProProValProAspHisSerAsp	110
540	CTGTGCAACAAGTACCGTGTGGGCCACGTGGATGTGCCACCTGTCCCCGACCACTCTGAC	481
150	LysGluAlaPheGlnArgLysLysLeuGlyCysGlnLeuLeuGlyThrTyrLysGlnVal	130
600	AAAGAAGCCTTCCAAAGGAAAAAGTTGGGTTGCCAGCTTCTGGGGACATACAAGCAAG	541
159	IleSerValValValGlnAlaPhe***	150
660	ATAAGTGTGGTGCCAGGCCTTCTAGAGAGGAGGTCTTGAATGTACCATGGACTGAGGG	601
720	ACCTCAGGAGCAGGATCCGGAGGTGGGGAGGGGGCTCAAAATGTGCTGGGGTTTGGGACA	661
780	TTGTTAAATGCAAAACGGGGCTGCTGGCAGACCCCAGGGATTTCCAGGTACTCACTGCAC	721
840	TCTGGGCTGGGCCATGATGGAATCTGGCAAAGTTGAAACTTCCATAGGCAGAGCTTCTAT	781
900	ACAGCCCAGCACCAGCTAGAAATGGCAATGAGGGTGTTGGTCTGAGAGATTTCTGTCTCA	841
	CTCACTCACTCACTCACTCACTCACT	901

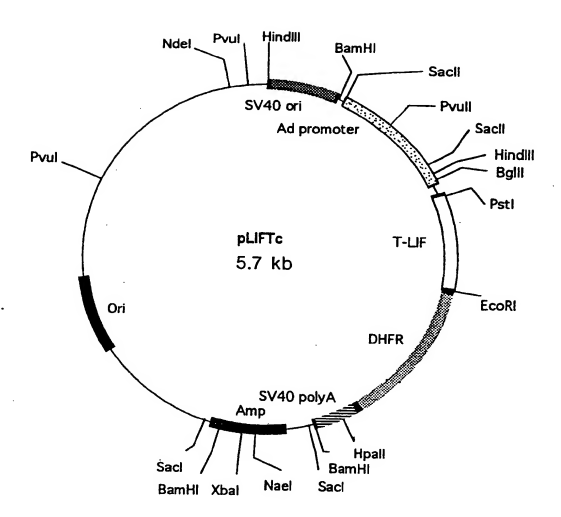
FIG. 26

60	GACCTTTTGCCTTTTCTCTCTCCTGGTGCACCATTTCCTCTCCCTCC	1
120	GTGCCCCTGCTGTTGGTTCTGCACTGGAAACATGGGGCGGGGAGCCCCCTCCCCATCACC	61
4	MetAsnGlnIle	1
180	CCTGTCAACGCCACCTGTGCCATACGCCACCCATGTCACAACAACCTCATGAACCAGATC	121
_ 24	ArgSerGlnLeuAlaGlnLeuAsnGlySerAlaAsnAlaLeuPheIleLeuTyrTyrThr	4
240	AGGAGCCAACTGGCACAGCTCAATGGCAGTGCCAATGCCCTCTTTATTCTCTATTACACA	181
44	AlaGlnGlyGluProPheProAsnAsnLeuAspLysLeuCysGlyProAsnValThrAsp	24
300	GCCCAGGGGGAGCCGTTCCCCAACAACCTGGACAAGCTATGTGGCCCCAACGTGACGGAC	241
64	PheProProPheHisAlaAsnGlyThrGluLysAlaLysLeuValGluLeuTyrArgIle	44
360	. TTCCCGCCCTTCCACGCCAACGGCACGGAGAAGGCCAAGCTGGTGGAGCTGTACCGCATA	301
84	ValValTyrLeuGlyThrSerLeuGlyAsnIleThrArgAspGlnLysIleLeuAsnPro	64
420	GTCGTGTACCTTGGCACCTCCCTGGGCAACATCACCCGGGACCAGAAGATCCTCAACCCC	361
104	SerAlaLeuSerLeuHisSerLysLeuAsnAlaThrAlaAspIleLeuArgGlyLeuLeu	84
480	AGTGCCCTCAGCCTCCACAGCAAGCTCAACGCCACCGCCGACATCCTGCGAGGCCTCCTT	421
124	SerAsnValLeuCysArgLeuCysSerLysTyrHisValGlyHisValAspValThrTyr	104
540	AGCAACGTGCTGTGCCGCCTGTGCAGCAAGTACCACGTGGGCCATGTGGACGTGACCTAC	481
144	${\tt GlyProAspThrSerGlyLysAspValPheGlnLysLysLeuGlyCysGlnLeuLeu}$	124
600	GGCCCTGACACCTCGGGTAAGGATGTCTTCCAGAAGAAGAAGCTGGGCTGTCAACTCCTG	541
159	GlyLysTyrLysGlnIleIleAlaValLeuAlaGlnAlaPhe***	144
660	GGGAAGTATAAGCAGATCATCGCCGTGTTGGCCCAGGCCTTCTAGCAGGAGGTCTTGAAG	601
720	TGTGCTGTGAACCGAGGGATCTCAGGAGTTGGGTCCAGATGTGGGGGCCTGTCCAAGGGT	661
	GGCTGGGCCCAGGGCATCGCTAAACCCAAATGGGGGCTGCTGGCTG	
840	TGGCCAGTCCACTCTGGGCTGGGCTGTGATGAAGCTGAGCAGAGTGGAAACTTCC	781
900	ATAGGGAGGAGCTAGAAGAAGGTGCCCCTTCCTCTGGGAGATTGTGGACTGGGAGCGT	841
960	GGGCTGGACTTCTGCCTCTACTTGTCCCTTTGGCCCCCTTGCTCACTTTGTGCAGTGAACA	901
	AACTACACAAGTCATCTACAAGAGCCCTGACC	961

FIG. 27



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Dihydrofolate reductase 3' end

Adenovirus promoter

SV40 origin of replication

SV40 PolyA signal

T-LIF coding region

Bacterial origin of replication

FIG. 29

A. Int. Cl. ⁶ C1	CLASSIFICATION OF SUBJECT MATTER 2N 15/54, 15/19, A61K 31/70, 35/16						
According to International Patent Classification (IPC) or to both national classification and IPC							
В.	FIELDS SEARCHED						
Minimum doo Electronic d	Minimum documentation searched (classification system followed by classification symbols) Electronic databases: WPAT, CASM. Both through QUESTEL ORBIT. Keywords as below.						
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic databases: BIOT, USPM, JAPIO, STN, Medline Embase. Keywords as below. AU IPC C12N 15/54, 15/19.							
1	ta base consulted during the international search (
Sensitiv:; Tr STN Sequen 1. AATGTO 2. YYTAO	CAA[AG]GGAA[GA]AGT[GA][GA]T; GCC GEPFPNNVEKLCAPNM;LGTSLGNITRDC DT MEDLINE, EMBASE (through DIALOG)	lant:; Graft:; Reduc:; Deplete#; Less:; Re r CASM only) CATTTTTGG[GA]GGAAC[AG]CC[CT]	eject:; Hyperacut:;				
C.	DOCUMENTS CONSIDERED TO BE RELEV	ANT					
Category*	Citation of document, with indication, where	appropriate, of the relevant passages	Relevant to Claim No.				
х	 M.S. Sandrin et al: "Anti-pig IgM antibodies in human serum react predominantly with Gal (α1-3) Gal epitopes". Proc Natl. Acad. Sci. USA, vol 90, pp 11391-11395, December 1993. See abstract; p 11393: section titled "Human Anti-pig Antibodies React Predominantly with Terminal Galactose Residues"; p 11393, column 2 last line to p 11394 column 2 line 2; p 11394: "Discussion" first paragraph; p 11395, first 2 sentences of 2nd full paragraph and first 10 lines 10 						
	paragraph; p 11395, first 2 sentences of 2nd last paragraph.	d full paragraph and first 10 lines 10					
	er documents are listed continuation of Box C.	X See patent family annex	.				
"A" docum not co earlier interna docum co whi anothe	all categories of cited documents: then defining the general state of the art which is insidered to be of particular relevance. It is document but published on or after the stional filling date in the publication of the stablish the publication date of a citation or other special reason (as specified) then the publication of other means then the prior to the international filing date or than the priority date claimed	"Y" document is taken along document of particular invention cannot be con inventive step when the with one or more other	ate and not in conflict . cited to understand the erlying the invention relevance; the claimed sidered novel or cannot be n inventive step when the erlevance; the claimed sidered to involve an document is combined such documents, such ous to a person skilled in				
	nual completion of the international search	Date of mailing of the international search	report				
20 June 1995		27 JUNE 1995 (27.06.	95)				
	iling address of the ISA/AU N INDUSTRIAL PROPERTY ORGANISATION T 2606	Authorized officer ROBYN PORTER) (·				
Facsimile No.	06 2853929	Telephone No. (06) 2832318					

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BNSDOCID: <WO_____9520661A1_I_>

Category *	Citation of document, with indication, where appropriate of the relevant passages	Relevant to Claim No.
P,X	WO 94/21799 (AUSTIN RESEARCH INSTITUTE) 29 September 1994. See entire specification	1-4, 8, 38, 39
X	M.S. Sandrin & I.F.C. McKenzie: "Galα(1,3)Gal, the Major Xenoantigen(s) Recognised in Pigs by Human Natural Antibodies". Immunological Reviews, 141, 1, pp 169-190, 1994. See page 174, last paragraph, lines 5-8; p 175 lines 4-5; p 176 lines 4-14; p 176, 2nd full paragraph; p 176, line 4 of last paragraph to end of paragraph; p 177, lines 20-30; p 178, Figure 2; p 184, Table III; p 186, lines 14-15 of "Summary".	1, 4-11, 38-43
P,X	WO 94/02616 (THE REGENTS OF THE UNIVERSITY OF MICHIGAN), 3 February 1995. See Sequence ID Numbers 13 and 14.	1
x	K. Gustafsson et al: " α 1,3Galactosyltransferase: A Target for in vivo Genetic Manipulation in Xenotransplantation".	1-3
Y	Immunological Reviews, 141, 1, pp 59-70, 1994. See p 63 lines 7-9; p 68 lines 9-11, 16-24.	4
Y	U. Galili: "Interaction of the natural anti-Gal antibody with α -galactosyl epitopes: a major obstacle for xenotransplantation in humans". Immunology Today, 14, 10, pp 480-482, 1993.	4, 8, 12-31
Y	R.D. Larsen et al: "Isolation of a cDNA encoding a murine UDPgalactose: β -D-galactosyl-1,4-N-acetyl-D-glucosaminide α -1,3-galactosyltransferase: Expressin cloning by gene transfer". Proc. Natl. Acad. Sci. USA, 86, pp 8227-8231, November 1989. See entire document but especially abstract; p 8227, column 2, last sentence of Introduction; p 8228, column 2 - "Results"; p 8231, last paragraph.	32-37
Y	F. A. Fletcher et al: "Leukemia Inhibitory Factory Improves Survival of Retroviral Vector-infected Hematopoietic Stem Cells In Vitro, Allowing Efficient Long-term Expression of Vector-encoded Human Adenosine Deaminase In Vivo. "The Journal of Experimental Medicine, 174, 4, pp 837-45, 1991. See Summary; Introduction; p 839, Section of Results titled "Effect of LIF on Tem Cell Survival In Vitro"; p 844, column 2, lines 10-12.	32-37
Y	WO 91/13985 (J. HEATH, A. SMITH & P. RATHJEN), 19 September 1991. See page 1 lines 3-18, page 2 lines 12-14, p 4 line 5 - p 5 line 1, p 5 lines 16-20, p 11 lines 21-26, Examples, Claims 1, 5, 6, 7.	32-37
P,Y	B. B. Samal et al: "High level expression of human leukemia inhibitory factor (LIF) from a synthetic gene in <u>Escherichia coli</u> and the physical and biological characterization of the protein". Biochimica et Biophysica Acta, 1260, pp 27-34, 1995. See entire document, especially Figures 2 and 3.	32-37
Y	T.A. Willson et al: "Cross-species comparison of the sequence of the leukaemia inhibitory factor gene and its protein". European Journal of Biochemistry, 204, 1, pp 21-30, 1992. See entire document, especially p 25 lines 36-39 and 41-57.	32-37
	N.M. Gough et al: "Molecular Cloning and expression of the human homologue of the murine gene encoding myeloid leukemia-inhibitory factor". Proc. Natl Acad Sci, 85, pp 2623-2627, April 1988. See entire document especially Figures 3 and 4.	32-37
J		

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BNSDOCID: <WO_____9520661A1_I_>

PCT/IB 95/00088

ategory *	Citation of document with in the	T
acegory	Citation of document, with indication, where appropriate of the relevant passages	Relevant to Claim No.
Y	D.P. Gearing et al: "Complete sequence of murine myeloid leukaemia inhibitory factor "(LIF)". Nucleic Acids Research, 16, 20, p 9857, 1988. See entire document.	32-37
Y	J.F. Moreau et al: "Leukaemia inhibitory factor is identical to the myeloid growth factor human interleukin for DA cells". Nature, 336, pp 690-692, 15 December 1988. See abstract, Figure 1.	32-37
Y	WO 88/07548 (AMRAD CORPORATION LIMITED) 6 October 1988. See page 1 lines 2-5, claims 1-3, 7, 8, 13-17, 34, 35.	32-37
	D.P. Gearing et al: "Production of Leukemia Inhibitory Factor in <u>Escherichia coli</u> by a Novel Procedure and Its Use in Maintaining Embryonic Stem Cells in Culture". Bio/Technology, 7, pp 1157-1161, November 1989. See abstract, p 1157, column 2, lines 2-10, 42-45, p 1159, column 2, lines 15 - end of paragraph.	32-37
	T. Yamamori et al: "The Cholinergic Neuronal Differentiation Factor from Heart Cells Is Identical to Leukemia Inhibitory Factor". Science, 246, pp 1412-1416, 15 December 1989. See entire document.	32-37
	D.G. Lowe et al: "Genomic Cloning and Heterologous Expression of Human Differentation-Stimulating Factor". DNA, 8, 5, pp 351-359, 1989. See whole document especially abstract; p 352, column 1, lines 1-8, 21-22; p 352 "Materials and Methods".	32-37
1 4	J. Stahl et al: "Structural Organization of the Genes for Murine and Human Leukemia Inhibitory Factor", Journal Biol Chem, 265, 15, pp 8833-41, 1990. See entire document.	32-37
Î	H.A. Vaughan et al: "Galα(1,3)Gal is the major xenoepitope expressed on pig endothelial cells recognized by naturally occurring cytoxic human antibodies". Transplantation, 58, 8, pp 879-882, 1994. See p 879, column 2 last 5 lines to p 880, column 1, last 5 lines of 'Materials and Methods'; p 880, column 1, last 10 lines 10 column 2, end of section; p 882, column 1, 11th line from bottom to end.	38-39
1 4	WO 93/16729 (BIOTRANSPLANT, INC) 2 September 1993. See page 3 lines 1-14 and 23-33; page 6 last 4 lines - page 7 line 2; page 14, section titled "IgM Depleting Technique"; claims 1, 8, 10.	40-41
x v	WO 92/07581 (AUTOIMMUNE, INC) 14 May 1992. See claim 11.	40
1 li li p	D. Latinne et al: "Depletion of IgM xenoreactive Natural Antibodies by Injection of anti-μ Monoclonal Antibodies". Immunological Reviews, 141, 1, pp 94-125, 1994. See page 98, full paragraph; page 99 lines 11-13; page 102, new section, ines 3-12 and 14-16; page 106 lines 3-11; page 110, sentence beginning on 3rd last ine to page 111, line 2; page 115, lines 3-11; page 116 sentence on last line age 117 line 1, page 117, lines 3-21 of "Conclusion"; page 118, paragraph 2 lines 1-5.	40-43
P	page 117, line 2; page 115, lines 3-11; page 116 sentence on last line - lage 117 line 1, page 117, lines 3-21 of "Conclusion"; page 118, paragraph 2 lines	·

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Information on patent family members

PCT/IB 95/00088

This Annex lists the known "A" publication level patent family members relating to the patent documents cited in the above-mentioned international search report. The Australian Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

	Patent Document Cited in Search Report				Patent Family	Member		
wo	9421799	·· ··· AU · ·	62792/94	32				-
wo	9402616	EP	654082	IL	106416	US	5324663	
wo	9113985	EP	518933	GB	9004890	JP	5504955	
wo	8807548	AT EP HU NO ZA	102991 285448 51330 950509 8802277	AU FI IL NZ US	15907/88 894613 85961 224105 5187077	DK HK NO PT	4831/89 336/95 885339 87133	
wo	9316729	AU	37796/93		·			
wo	9207581	AU EP NO	89426/91 555413 931536	BR HU	9107055 63957	CA IL	2093513 99864	
wo	9008188	AU NZ	48356/90 232072	CA US	2045126 5418159	EP	453453	
EP	235805	AU FI ZA	69691/87 870916 8701562	DK JP	1098/87 62281824	EP NO	235805 870866	

END OF ANNEX

Box II (continued)

The inventors have established that most xenoantibodies are directed to a terminal α 1,3 galactose linkage (termed the GAL epitope) residue on the cell surface of graft tissues.

Using this observation they have developed several strategies for limiting graft rejection.

Invention 1 as defined in claims 1 to 31 is to eliminate the GAL epitope on the donor organ (as described on page 15) by interfering with the expression of the enzyme responsible for forming the α 1,3 galactose linkage.

Invention 2 as defined in claims 38 to 39 is to block the circulating GAL antibodies in the recipient by IV administration of an α 1,3 galactose which will bind to the antibody. Other sugars can also block the antibody when administered intravenously (see pages 21 to 22).

Invention 3 as defined in claims 40 to 45 is to non-specifically deplete the recipient of total IgM prior to transplantation t reduce the acute phase of Ig response (see page 22).

Invention 4 as defined in claims 32 to 37 is directed to a novel LIF. LIF has been identified as a suitable differentiation inhibiting factor for growth of ES cells having an inactivated α 1,3 GalT gene for the development of a transgenic animal (see page 30).

Since inventions 2 and 3 are directed at either blocking or depleting the recipient's Ig levels and invention 1 is directed at eliminating the GAL epitope on the donor organs, the international application does not comply with the requirements of unity of invention because the inventions defined do not share a 'technical relationship' and thus these inventions do not relate to one invention or to a single inventive concept.

Further, since invention 1 is directed at eliminating the GAL epitope on the donor organs and invention 4 is directed to an LIF, there is also no 'technical relationship' between these 2 inventions and consequently the international application does not elate to one invention or to a single inventive concept.

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	Box I Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)
	This international search report has not established in respect of certain claims under Article 17(2)(a) for the following reasons:
	Claims Nos.: 18, 26 and 29 because they relate to subject matter not required to be searched by this Authority, namely:
.	They include humans within their scope. However, the claims were searched as if they specifically excluded humans ie " non-human mammals".
	Claim Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
	Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
	This International Searching Authority found multiple inventions in this international application, as follows:
	Claims 1 to 31; Claims 32 to 37; Claims 38 to 39; and Claims 40 to 45
	As reasoned on "extra" sheet.
	As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims
	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
	As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
	No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
	Remark on Protest
	The additional search fees were accompanied by the applicant's protest.
	X No protest accompanied the payment of additional search fees.

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BNSDOCID: <WO_____9520661A1_I_>

C(Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT						
Category *	Citation of document, with indication, where appropriate of the relevant passages	Relevant to Claim No.				
P,Y	M. Soares et al: "In Vivo IgM depletion by Anti-μ Monoclonal Antibody Therapy". Transplantation, 57, 7, pp 1003-1009, 1994. See abstract lines 23-27; p 1003 column 2, lines 1-10 and 33-39; p 1004, column 1, lines 6-9; column 2 lines 3-5; p 1006, figure 3, column 2, lines 4-8, p 1007, column 2 lines 2-5 and 11-28; p 1009, column 1, lines 20-24.	40, 41				
х	M. Soares et al: "In Vivo Depletion of Xenoreactive Antibodies with an Anti-μ Monoclonal Antibody". Transplantation, 56, 6, pp 1427-1433, 1993. See abstract, 2nd sentence, p 1428, column 1, last full sentence, p 1429, paragraph spanning columns 1 and 2, Figure 1; p 1429, column 2, lines 16-43, p 1430, figures 3 and 4; p 1431, 1st sentence of 'DISCUSSION'.	40,41				
X	L. Gambiez et al: "The Role of Natural IgM in the Hyperacute Rejection of Discordant Heart Xenografts". Transplantation, 54, 4, pp 577-583, 1992. See last paragraph of abstract; p 577, column 2, last sentence of introduction; p 579, column 1 line 19 - column 2 line 14; p 580, column 1 line 22-23, 29-30 and column 2 lines 14-20; p 582, column 1, last full sentence.	40, 41				
P,A	R. Oriol et al: "Monomorphic and polymorphic carbohydrate antigens on pig tissues: implications for organ xenotransplantation in the pig-to-human model". Transplant Internationa, 7, 6, pp 405-413, 1994.	1-7, 12, 15-18, 21-26 28, 29, 31				
A	WO 90/08188 (AMRAD CORPORATION LIMITED) 26 July 1990.	32-37				
A	EP 235805 (THE ROYAL FREE HOSPITAL SCHOOL OF MEDICINE) 9 September 1987.					
A	G. Hale et al: "Removal of T Cells From Bone Marrow for Transplanation: A Monoclonal Antilymphocyte Antibody That Fixes Human Complement". Blood, 62, 4, pp 873-882, 1983.					
	·					

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